

**JUNE 2014 GROUNDWATER SAMPLING DATA  
SUMMARY REPORT  
BETHPAGE, NY**

**Prepared for:**



**Department of the Navy  
Naval Facilities Engineering Command, Mid-Atlantic  
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**CTO WE15**

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### **List of Acronyms and Abbreviations**

DOT	Department of Transportation
IDW	Investigation Derived Waste
Katahdin	Katahdin Analytical Services, Inc.
NWIRP	Naval Weapons Industrial Reserve Plant
ONCT	Onsite Containment System
OU	Operable Unit
POTW	Publicly Owned Treatment Works
QA	Quality Assurance
QC	Quality Control
SAP	Sampling and Analysis Plan
UFP	Uniform Federal Policy
VOC	Volatile Organic Compounds

## **1. PROJECT BACKGROUND**

Resolution Consultants has prepared this Data Summary Report for the Naval Facilities Engineering Command, Mid-Atlantic under contract task order WE15 Contract N62470-11-D-8013. The report describes monitoring well sampling activities in June 2014 for the Naval Weapons Industrial Reserve Plant (NWIRP) Bethpage Operable Unit (OU) 2 Site 1 offsite plume. NWIRP Bethpage is located in east-central Nassau County, Long Island, New York, approximately 30 miles east of New York City (Figure 1).

This data summary report provides information on sampling 13 monitoring wells. The purpose of this sampling is to provide information on the extent and magnitude of volatile organic compounds (VOCs) located in a narrow area immediately south of the Onsite Containment System (ONCT) in the western offsite plume, which could represent contamination that has bypassed the ONCT. The locations of monitoring wells sampled as part of this effort are shown in Figure 2.

Documentation of these activities is included in the appendices of this report. Appendix A contains the groundwater sampling forms, Appendix B contains analytical lab sheets, and Appendix C contains documentation of data validation.

## **2. FIELD PROGRAM**

Field tasks were conducted in June of 2014 in accordance with the Uniform Federal Policy (UFP) Sampling and Analysis Plan (SAP) Addendum: Groundwater Sampling Using Low Stress (Low Flow) Purging and Sampling Protocol (Resolution Consultants, 2013). The field investigation included purging and sampling of the 13 monitoring wells.

### **2.1 Sampling**

Wells were purged with a submersible pump with the intake placed at the approximate midpoint of the screened interval. The following field water quality parameters were continuously measured during purging: water temperature, pH, conductivity, oxidation-reduction potential, dissolved oxygen and turbidity. Groundwater analytical samples were collected when field water quality parameters stabilized. Samples were analyzed for VOCs via Method 8260B and 1,4-dioxane via Method 8270C by Katahdin Analytical Services (Katahdin). All purge water was managed as investigation derived waste (IDW). Samples were placed in a cooler containing ice and held for sample pick up by the laboratory courier. All samples were submitted to the laboratory for analyses of VOCs for the analytes listed in, and in accordance with, GC method SW846-8260B. Quality assurance (QA) and quality control (QC) samples were collected during the sampling effort.

Resolution Consultants utilized dedicated and disposable sampling equipment when possible to avoid the potential for cross-impacting of samples. The sampling equipment included dedicated disposable polyethylene tubing, disposable gloves, and laboratory supplied sample bottles. Hand held equipment was decontaminated using an alconox and water wash, a potable water rinse followed by a distilled water rinse. Water was collected in 5-gallon pails or 55-gallon drums.

### **2.2 Investigation Derived Waste**

Purge water was transported from point of generation to the designated staging area at NWIRP in Department of Transportation (DOT) approved 55-gallon drums. Purge water was then containerized in a frac tank and stored at NWIRP Bethpage for characterization and ultimate disposal to the Nassau County Publicly Owned Treatment Works (POTW) in accordance with the facility's existing discharge permit. A representative water sample will be collected from each of the frac tanks and submitted to Katahdin for analysis. No solid waste was generated during sampling.

### 3. **SUMMARY**

Well construction information is summarized in Table 1; analytical data is summarized in Table 2; stabilized field water quality parameters are summarized in Table 3. Groundwater sample logs, lab analytical sheets, and data validation packages are included in Appendix A, B and C, respectively.

#### **4. REFERENCES**

Resolution Consultants, 2013. UFP SAP Addendum, *Groundwater Sampling Using Low Stress (Low Flow) Purging and Sampling* Protocol. November.

## Tables

Table 1.  
 Monitoring Well  
 Construction Summary

Well	Total Depth (ft bgs)	Top of Screen (ft bgs)	Bottom of Screen (ft bgs)	Mid-screen (ft bgs)	Sump Length (ft)	VPB affiliation
RE103 D1	645	625	640	630	5	VPB137
RE103 D2	673	653	673	663	0	
RE103 D3	735	715	730	720	5	
RE104 D1	375	350	370	360	5	VPB138
RE104 D2	735	710	730	720	5	
RE104 D3	785	760	780	770	5	
RE105 D1	554.9	530	550	540	5	VPB139
RE105 D2	755.9	730	750	740	5	
RE108D1	545	530	550	540	5	VPB142
RE108D2	655	630	650	640	5	
BPOW5-1	515	480	510	495	5	VPB132
BPOW5-2	585	540	580	560	5	
BPOW5-3	665	620	660	640	5	

Table 2. Analytical Data Summary

Location	NYSDEC	RE103D1	RE103D2	RE103D3	RE104D1
Sample Date	Groundwater	6/11/2014	6/11/2014	6/11/2014	6/12/2014
Sample ID	Guidance or	RE103D1-GW-061114	RE103D2-GW-061114	RE103D3-GW-061114	RE104D1-GW-061214
Sample type code	Standard Value	N	N	N	N
(Note 1)	(Note 1)				
<b>VOC 8260B (ug/L)</b>					
1,1,1-TRICHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	<b>0.41 J</b>
1,1,2,2-TETRACHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1,2-TRICHLORO-1,2,2-TRIFLUOROETHANE	5	<b>18</b>	<b>3.8</b>	<b>2.9</b>	<b>6.7</b>
1,1,2-TRICHLOROETHANE	1	<b>0.68 J</b>	<b>0.37 J</b>	< 0.50 U	< 0.50 U
1,1-DICHLOROETHANE	5	<b>1.3</b>	<b>0.67 J</b>	<b>0.64 J</b>	<b>0.44 J</b>
1,1-DICHLOROETHENE	5	<b>9.4</b>	<b>0.61 J</b>	<b>0.71 J</b>	<b>1.5</b>
1,2,4-TRICHLOROBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DIBROMO-3-CHLOROPROPANE	0.04	< 0.75 U	< 0.75 U	< 0.75 U	< 0.75 U
1,2-DIBROMOETHANE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROETHENE, TOTAL	5	<b>4.2</b>	<b>1.1 J</b>	<b>0.87 J</b>	<b>1.6 J</b>
1,2-DICHLOROPROPANE	1	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,3-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,4-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,4-DIOXANE	NL	<b>20</b>	<b>1.0</b>	<b>0.92</b>	<b>12</b>
2-BUTANONE	50	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ
2-HEXANONE	50	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ
4-METHYL-2-PENTANONE	NL	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ
ACETONE	50	< 2.5 U	< 2.5 U	< 2.5 U	< 2.5 U
BENZENE	1	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
BROMODICHLOROMETHANE	50	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
BROMOFORM	50	< 0.50 UJ	< 0.50 UJ	< 0.50 UJ	< 0.50 UJ
BROMOMETHANE	5	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
CARBON DISULFIDE	60	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CARBON TETRACHLORIDE	5	<b>0.54 J</b>	< 0.50 U	<b>0.30 J</b>	< 0.50 U
CHLOROBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CHLOROETHANE	5	< 1.0 UJ	< 1.0 UJ	< 1.0 UJ	< 1.0 UJ
CHLOROFORM	7	<b>0.95 J</b>	<b>0.96 J</b>	<b>0.94 J</b>	<b>0.37 J</b>
CHLOROMETHANE	5	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
CIS-1,2-DICHLOROETHENE	5	<b>4.2</b>	<b>1.1</b>	<b>0.87 J</b>	<b>1.6</b>
CIS-1,3-DICHLOROPROPENE	0.4	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CYCLOHEXANE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
DIBROMOCHLOROMETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
DICHLORODIFLUOROMETHANE	5	<b>0.54 J</b>	< 1.0 U	< 1.0 U	<b>0.98 J</b>
ETHYLBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
ISOPROPYLBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
M- AND P-XYLENE	NL	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
METHYL ACETATE	NL	< 0.75 U	< 0.75 U	< 0.75 U	< 0.75 U
METHYL CYCLOHEXANE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
METHYL TERT-BUTYL ETHER	10	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
METHYLENE CHLORIDE	5	< 2.5 U	< 2.5 U	< 2.5 U	< 2.5 U
O-XYLENE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
STYRENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TETRACHLOROETHENE	5	<b>4.9</b>	<b>0.77 J</b>	< 0.50 U	<b>2.4</b>
TOLUENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TRANS-1,2-DICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TRANS-1,3-DICHLOROPROPENE	0.4	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TRICHLOROETHENE	5	<b>1200 J</b>	<b>670 J</b>	<b>510 J</b>	<b>160</b>
TRICHLOROFUOROMETHANE	5	<b>0.25 J</b>	< 1.0 U	< 1.0 U	< 1.0 U
VINYL CHLORIDE	2	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
XYLENES, TOTAL	5	< 1.5 U	< 1.5 U	< 1.5 U	< 1.5 U

Table 2. Analytical Data Summary

Location	NYSDEC	RE104D2	RE104D3	RE105D1	RE105D1
Sample Date	Groundwater	6/12/2014	6/12/2014	6/11/2014	6/11/2014
Sample ID	Guidance or	RE104D2-GW-061214	RE104D3-GW-061214	RE105D1-GW-061114	GWDUP-061114
Sample type code	Standard Value	N	N	N	FD
(Note 1)	(Note 1)				
<b>VOC 8260B (ug/L)</b>					
1,1,1-TRICHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	<b>0.62 J</b>
1,1,2,2-TETRACHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1,2-TRICHLORO-1,2,2-TRIFLUOROETHANE	5	< 0.50 U	< 0.50 U	<b>12</b>	<b>13</b>
1,1,2-TRICHLOROETHANE	1	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1-DICHLOROETHANE	5	< 0.50 U	< 0.50 U	<b>0.39 J</b>	<b>0.40 J</b>
1,1-DICHLOROETHENE	5	< 0.50 U	< 0.50 U	<b>1.5</b>	<b>1.7</b>
1,2,4-TRICHLOROBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DIBROMO-3-CHLOROPROPANE	0.04	< 0.75 U	< 0.75 U	< 0.75 U	< 0.75 U
1,2-DIBROMOETHANE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROETHENE, TOTAL	5	<b>0.84 J</b>	< 1.0 U	<b>1.9 J</b>	<b>1.9 J</b>
1,2-DICHLOROPROPANE	1	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,3-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,4-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,4-DIOXANE	NL	< 0.17 U	< 0.17 U	<b>15</b>	<b>18</b>
2-BUTANONE	50	< 2.5 U	< 2.5 U	< 2.5 UJ	< 2.5 UJ
2-HEXANONE	50	< 2.5 U	< 2.5 U	< 2.5 UJ	< 2.5 UJ
4-METHYL-2-PENTANONE	NL	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ
ACETONE	50	< 2.5 U	< 2.5 U	< 2.5 U	<b>6.8 J</b>
BENZENE	1	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
BROMODICHLOROMETHANE	50	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
BROMOFORM	50	< 0.50 UJ	< 0.50 UJ	< 0.50 UJ	< 0.50 UJ
BROMOMETHANE	5	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
CARBON DISULFIDE	60	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CARBON TETRACHLORIDE	5	< 0.50 UJ	< 0.50 UJ	<b>0.24 J</b>	<b>0.27 J</b>
CHLOROENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CHLOROETHANE	5	< 1.0 UJ	< 1.0 UJ	< 1.0 UJ	< 1.0 UJ
CHLOROFORM	7	< 0.50 U	< 0.50 U	<b>0.50 J</b>	<b>0.47 J</b>
CHLOROMETHANE	5	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
CIS-1,2-DICHLOROETHENE	5	<b>0.84 J</b>	< 0.50 U	<b>1.9</b>	<b>1.9</b>
CIS-1,3-DICHLOROPROPENE	0.4	< 0.50 UJ	< 0.50 UJ	< 0.50 U	< 0.50 U
CYCLOHEXANE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
DIBROMOCHLOROMETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
DICHLORODIFLUOROMETHANE	5	< 1.0 UJ	< 1.0 UJ	<b>1.4 J</b>	<b>1.4 J</b>
ETHYLBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
ISOPROPYLBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
M- AND P-XYLENE	NL	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
METHYL ACETATE	NL	< 0.75 U	< 0.75 U	< 0.75 U	< 0.75 U
METHYL CYCLOHEXANE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
METHYL TERT-BUTYL ETHER	10	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
METHYLENE CHLORIDE	5	< 2.5 U	< 2.5 U	< 2.5 U	< 2.5 U
O-XYLENE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
STYRENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TETRACHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TOLUENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TRANS-1,2-DICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TRANS-1,3-DICHLOROPROPENE	0.4	< 0.50 UJ	< 0.50 UJ	< 0.50 U	< 0.50 U
TRICHLOROETHENE	5	<b>1.8 J</b>	< 0.50 U	<b>130</b>	<b>140</b>
TRICHLOROFUOROMETHANE	5	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
VINYL CHLORIDE	2	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
XYLENES, TOTAL	5	< 1.5 U	< 1.5 U	< 1.5 U	< 1.5 U

Table 2. Analytical Data Summary

Location	NYSDEC	RE105D2	RE108D1	RE108D2	BPOW5-1
Sample Date	Groundwater	6/11/2014	6/10/2014	6/10/2014	6/13/2014
Sample ID	Guidance or	RE105D2-GW-061114	RE108D1-GW-061014	RE108D2-GW-061014	BPOW5-1-GW-061314
Sample type code	Standard Value	N	N	N	N
(Note 1)	(Note 1)				
<b>VOC 8260B (ug/L)</b>					
1,1,1-TRICHLOROETHANE	5	0.71 J	< 0.50 U	1.3	< 0.50 U
1,1,2,2-TETRACHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1,2-TRICHLORO-1,2,2-TRIFLUOROETHANE	5	32	0.58 J	8.7	< 0.50 U
1,1,2-TRICHLOROETHANE	1	1.2	< 0.50 U	1.8	< 0.50 U
1,1-DICHLOROETHANE	5	1.5	< 0.50 U	5.8	< 0.50 U
1,1-DICHLOROETHENE	5	6.2	< 0.50 U	8.2	< 0.50 U
1,2,4-TRICHLOROBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DIBROMO-3-CHLOROPROPANE	0.04	< 0.75 U	< 0.75 U	< 0.75 U	< 0.75 U
1,2-DIBROMOETHANE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-DICHLOROETHENE, TOTAL	5	3.1	< 1.0 U	9.9	< 1.0 U
1,2-DICHLOROPROPANE	1	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,3-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,4-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,4-DIOXANE	NL	6.2	7.1	6.9	< 0.17 U
2-BUTANONE	50	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ
2-HEXANONE	50	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ	< 2.5 U
4-METHYL-2-PENTANONE	NL	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ	< 2.5 UJ
ACETONE	50	< 2.5 U	< 2.5 U	< 2.5 U	< 2.5 U
BENZENE	1	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
BROMODICHLOROMETHANE	50	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
BROMOFORM	50	< 0.50 UJ	< 0.50 UJ	< 0.50 UJ	< 0.50 U
BROMOMETHANE	5	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
CARBON DISULFIDE	60	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CARBON TETRACHLORIDE	5	4.7	< 0.50 U	2.0	< 0.50 U
CHLOROBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CHLOROETHANE	5	< 1.0 UJ	< 1.0 UJ	< 1.0 UJ	< 1.0 UJ
CHLOROFORM	7	2.3	< 0.50 U	4.2	< 0.50 U
CHLOROMETHANE	5	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
CIS-1,2-DICHLOROETHENE	5	3.1	< 0.50 U	9.9	< 0.50 U
CIS-1,3-DICHLOROPROPENE	0.4	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
CYCLOHEXANE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
DIBROMOCHLOROMETHANE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
DICHLORODIFLUOROMETHANE	5	0.58 J	< 1.0 U	0.27 J	< 1.0 UJ
ETHYLBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
ISOPROPYLBENZENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
M- AND P-XYLENE	NL	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
METHYL ACETATE	NL	< 0.75 U	< 0.75 U	< 0.75 U	< 0.75 U
METHYL CYCLOHEXANE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
METHYL TERT-BUTYL ETHER	10	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
METHYLENE CHLORIDE	5	< 2.5 U	< 2.5 U	< 2.5 U	< 2.5 U
O-XYLENE	NL	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
STYRENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TETRACHLOROETHENE	5	0.77 J	0.59 J	1.6	< 0.50 U
TOLUENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TRANS-1,2-DICHLOROETHENE	5	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TRANS-1,3-DICHLOROPROPENE	0.4	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
TRICHLOROETHENE	5	1500 J	82	3400 J	< 0.50 U
TRICHLOROFUOROMETHANE	5	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
VINYL CHLORIDE	2	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
XYLENES, TOTAL	5	< 1.5 U	< 1.5 U	< 1.5 U	< 1.5 U

Table 2. Analytical Data Summary

Location	NYSDEC	BPOW5-2	BPOW5-3
Sample Date	Groundwater	6/13/2014	6/13/2014
Sample ID	Guidance or	BPOW5-2-GW-061314	BPOW5-3-GW-061314
Sample type code	Standard Value	N	N
(Note 1)			
VOC 8260B (ug/L)			
1,1,1-TRICHLOROETHANE	5	< 0.50 U	< 0.50 U
1,1,2-TETRACHLOROETHANE	5	< 0.50 U	< 0.50 U
1,1,2-TRICHLORO-1,2,2-TRIFLUOROETHANE	5	< 0.50 U	< 0.50 U
1,1,2-TRICHLOROETHANE	1	< 0.50 U	< 0.50 U
1,1-DICHLOROETHANE	5	< 0.50 U	< 0.50 U
1,1-DICHLOROETHENE	5	< 0.50 U	< 0.50 U
1,2,4-TRICHLOROBENZENE	5	< 0.50 U	< 0.50 U
1,2-DIBROMO-3-CHLOROPROPANE	0.04	< 0.75 U	< 0.75 U
1,2-DIBROMOETHANE	NL	< 0.50 U	< 0.50 U
1,2-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U
1,2-DICHLOROETHANE	5	< 0.50 U	< 0.50 U
1,2-DICHLOROETHENE, TOTAL	5	< 1.0 U	< 1.0 U
1,2-DICHLOROPROPANE	1	< 0.50 U	< 0.50 U
1,3-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U
1,4-DICHLOROBENZENE	3	< 0.50 U	< 0.50 U
1,4-DIOXANE	NL	< 0.17 U	0.55
2-BUTANONE	50	< 2.5 UJ	< 2.5 UJ
2-HEXANONE	50	< 2.5 U	< 2.5 U
4-METHYL-2-PENTANONE	NL	< 2.5 UJ	< 2.5 UJ
ACETONE	50	< 2.5 U	< 2.5 U
BENZENE	1	< 0.50 U	< 0.50 U
BROMODICHLOROMETHANE	50	< 0.50 U	< 0.50 U
BROMOFORM	50	< 0.50 U	< 0.50 U
BROMOMETHANE	5	< 1.0 U	< 1.0 U
CARBON DISULFIDE	60	< 0.50 U	< 0.50 U
CARBON TETRACHLORIDE	5	< 0.50 U	< 0.50 U
CHLOROBENZENE	5	< 0.50 U	< 0.50 U
CHLOROETHANE	5	< 1.0 UJ	< 1.0 UJ
CHLOROFORM	7	< 0.50 U	< 0.50 U
CHLOROMETHANE	5	< 1.0 U	< 1.0 U
CIS-1,2-DICHLOROETHENE	5	< 0.50 U	< 0.50 U
CIS-1,3-DICHLOROPROPENE	0.4	< 0.50 U	< 0.50 U
CYCLOHEXANE	NL	< 0.50 U	< 0.50 U
DIBROMOCHLOROMETHANE	5	< 0.50 U	< 0.50 U
DICHLORODIFLUOROMETHANE	5	< 1.0 UJ	< 1.0 UJ
ETHYLBENZENE	5	< 0.50 U	< 0.50 U
ISOPROPYLBENZENE	5	< 0.50 U	< 0.50 U
M- AND P-XYLENE	NL	< 1.0 U	< 1.0 U
METHYL ACETATE	NL	< 0.75 U	< 0.75 U
METHYL CYCLOHEXANE	NL	< 0.50 U	< 0.50 U
METHYL TERT-BUTYL ETHER	10	< 0.50 U	< 0.50 U
METHYLENE CHLORIDE	5	< 2.5 U	< 2.5 U
O-XYLENE	NL	< 0.50 U	< 0.50 U
STYRENE	5	< 0.50 U	< 0.50 U
TETRACHLOROETHENE	5	< 0.50 U	< 0.50 U
TOLUENE	5	< 0.50 U	< 0.50 U
TRANS-1,2-DICHLOROETHENE	5	< 0.50 U	< 0.50 U
TRANS-1,3-DICHLOROPROPENE	0.4	< 0.50 U	< 0.50 U
TRICHLOROETHENE	5	< 0.50 U	< 0.50 U
TRICHLOROFLUOROMETHANE	5	< 1.0 U	< 1.0 U
VINYL CHLORIDE	2	< 1.0 U	< 1.0 U
XYLENES, TOTAL	5	< 1.5 U	< 1.5 U

**Notes:**

1 New York State Department of Environmental Conservation Division of Water Technical and Operation Guidance series  
(6 NYCRR 700-706, Part 703.5 summarized in TOGS 1.1.1)

Ambient water quality standards and groundwater effluent limitations, class GA; NL = Not Listed

**Bold** = Detected; **Bold and Italics** = Not detect exceeds NYS Groundwater Standards or guidance value

Yellow highlighted values exceed Groundwater Standards or guidance value

Sample type codes: N - normal environmental sample, FD - field duplicate

U = Nondetected result. The analyte was analyzed for, but was not detected above the reported sample quantitation limit.

UJ = The analyte was not detected above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte.

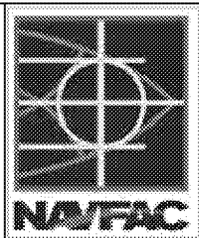
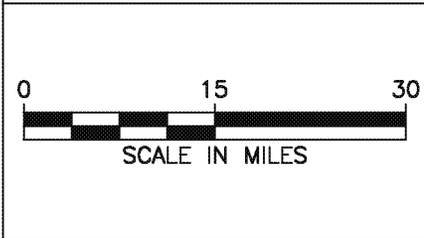
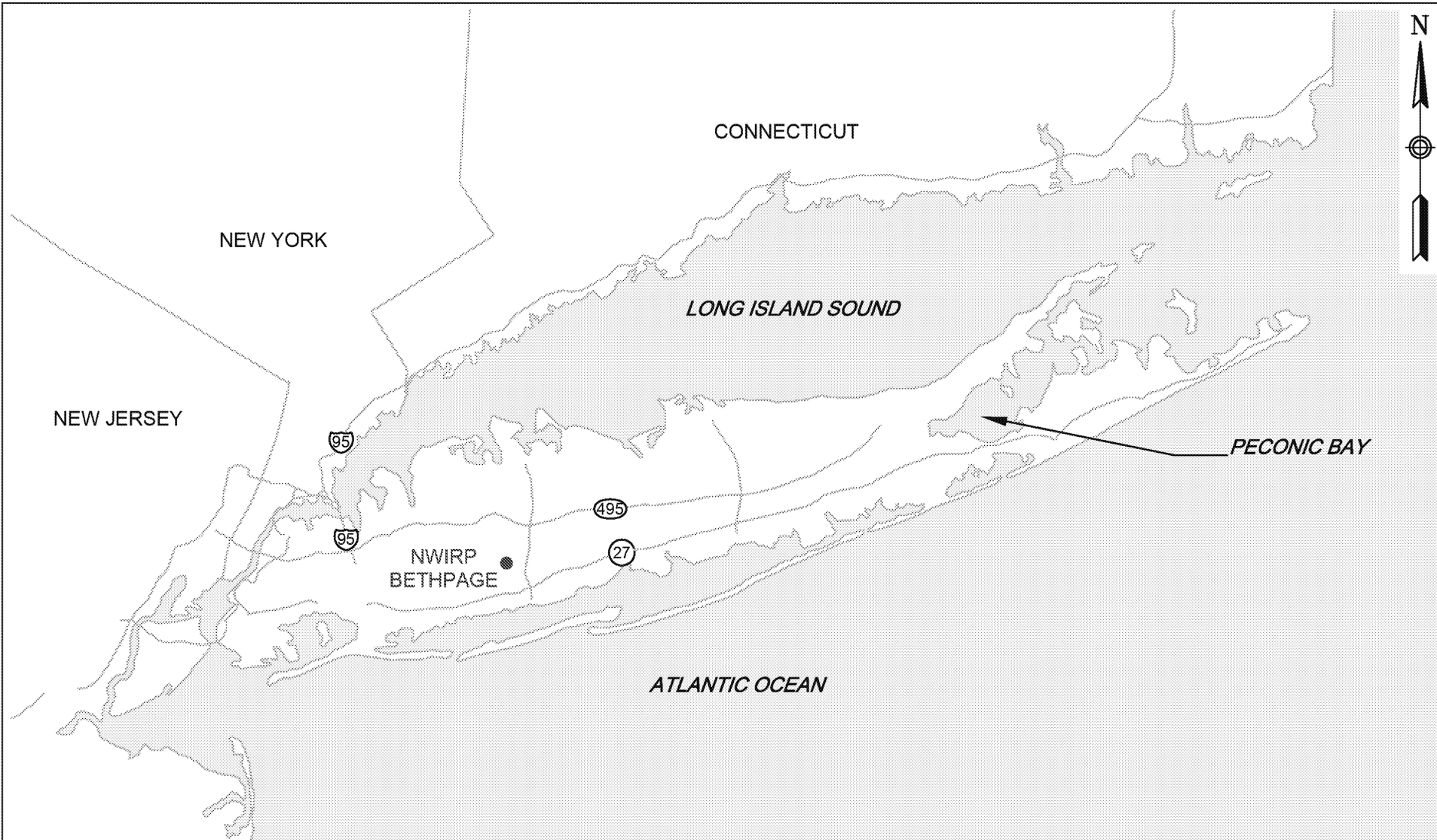
J = The analyte was positively identified; the associated numerical value is the approximate concentration of the analyte in the sample.

Table 3.  
 Stabilized Field Parameters

Well	Date	Temperature (°C)	pH	Specific Conductance (µS/cm)	DO (mg/L)	ORP (mV)	Turbidity (NTU)	Depth to water (ft bgs)	Flow rate (ml/min)	Drawdown (ft)
RE103 D1	6/11/2014	15.3	4.83	101	4.32	293.8	0.17	39.45	450	*
RE103 D2	6/11/2014	16.26	4.56	31	7.9	46.1	14.60	40.43	400	0
RE103 D3	6/11/2014	15.4	4.07	30	4.82	290.3	1.29	39.48	450	*
RE104 D1	6/12/2014	16.24	4.64	65	3.71	226.9	1.01	35.78	400	0
RE104 D2	6/12/2014	15.3	4.46	19	4.27	151.1	3.54	38.55	500	*
RE104 D3	6/12/2014	15.5	4.10	20	4.06	175.5	18.00	40.75	400	*
RE105 D1	6/11/2014	15.14	5.01	111	1.80	102.1	1.06	36.5	450	0
RE105 D2	6/11/2014	15.3	4.73	59	4.07	210.7	0.60	37.28	450	*
RE108 D1	6/10/2014	15.9	5.02	0.093	5.82	112.8	1.46	39.38	300	*
RE108 D2	6/10/2014	15.53	4.87	0.076	5.05	316.2	0.67	40.04	500	*
BPOW5-1	6/13/2014	20.08	4.29	96	0.67	91.7	21.20	20.92	450	0
BPOW5-2	6/13/2014	16.19	4.07	76	0.63	130.7	8.78	21.22	450	0
BPOW5-3	6/13/2014	15.9	4.55	30	0.45	148.4	8.59	21.34	550	*

\* Initial water level not equilibrated due to pump installation; drawdown during sampling not determined.

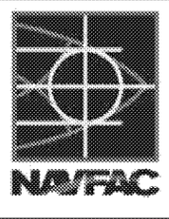
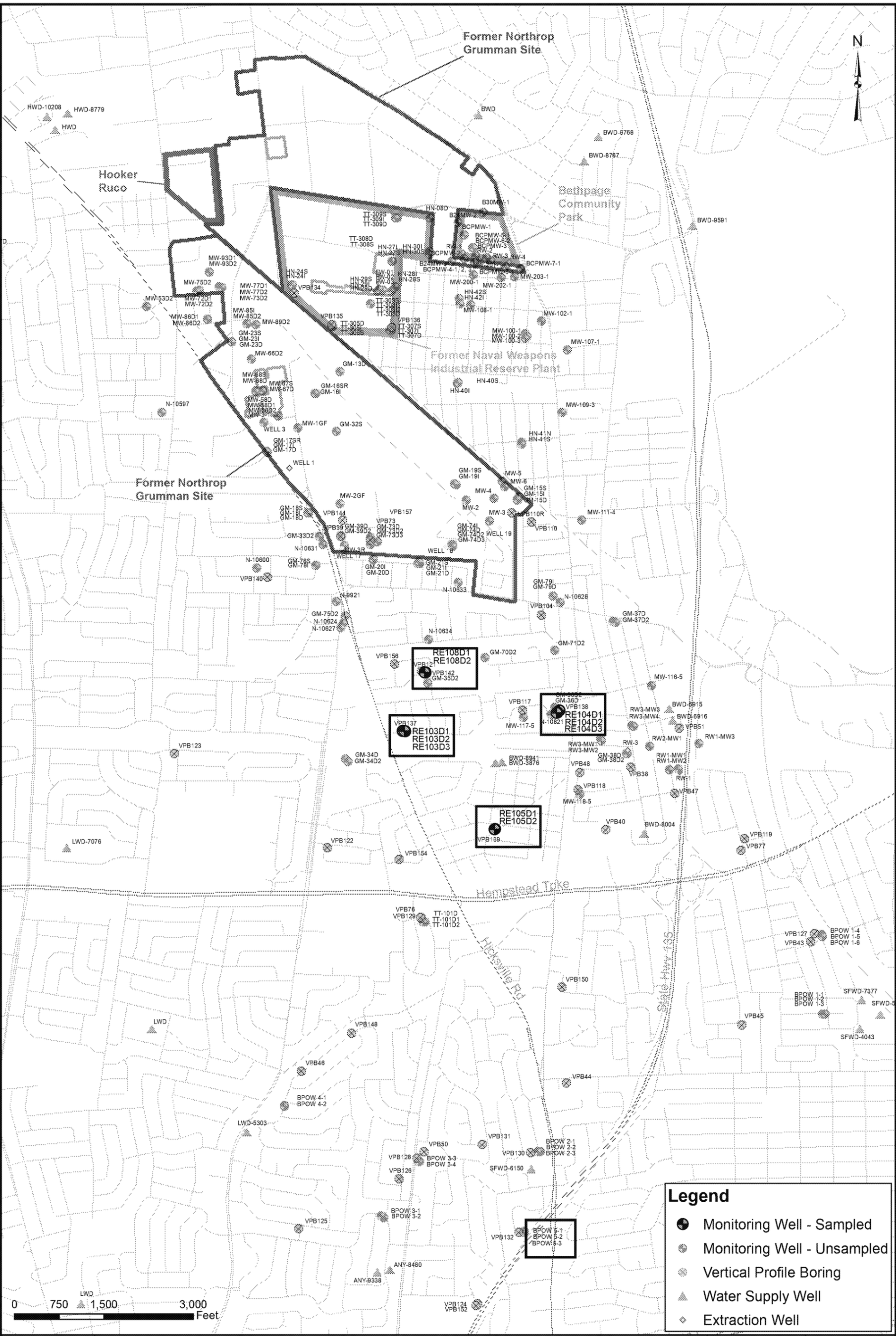
## Figures



GENERAL LOCATION MAP  
 NWIRP BETHPAGE  
 BETHPAGE, NEW YORK

CONTRACT NUMBER N62470-11-D-8013		CTO NUMBER WE15	
APPROVED BY ---		DATE ---	
APPROVED BY ---		DATE ---	
FIGURE NO. 1			REV 0

F:\Projects\Navy\Bethpage\WEXE087.0\_Deliverables\7.2\_CADD\GIS\_files\Bethpage\MAP\_DOCS\WIXD\MISC\2014\_03\_Wells\_Sampled\F2\_GW\_Samples\_2014\_06.mxd



**LOCATION MAP**  
**JUNE 2014 GROUNDWATER SAMPLING**  
**NAVAL WEAPONS INDUSTRIAL RESERVE PLANT**  
**BETHPAGE, NEW YORK**

Legend	
	Monitoring Well - Sampled
	Monitoring Well - Unsampled
	Vertical Profile Boring
	Water Supply Well
	Extraction Well

CONTRACT NUMBER N62470-11-D8013	CTO NUMBER WE15
APPROVED BY	DATE
APPROVED BY	DATE
FIGURE NO. 2	REV 0

## **Appendices**

**Appendix A**  
**Groundwater Sampling Forms**



RESOLUTION CONSULTANTS

# Low Flow Ground Water Sample Collection Record

Well ID: MW-10801

Client: Navy NMWRP Bethesda Date: 6/10/14 Time: Start 900 am/pm  
 Project No: 60266526 Finish 1230 am/pm  
 Site Location: ceiling of Corona  
 Weather Conds: cloudy, misting Collector(s): Pranath & Christopher

### 1. WATER LEVEL DATA: (measured from Top of Casing)

a. Total Well Length 545 c. Length of Water Column \_\_\_\_\_ (a-b) Casing Diameter/Material \_\_\_\_\_  
 b. Water Table Depth 40.10 d. Calculated System Volume (see back) 17gal 4" PVC

### 2. WELL PURGE DATA

a. Purge Method: Geotek bladder pump

b. Acceptance Criteria defined (see workplan)

- Temperature ± 3% - D.O. ± 10% (values >0.5 mg/L) Turbidity ± 10%
- pH ± 0.1 unit - ORP ± 10mV
- Sp. Cond. ± 3% - Drawdown < 0.3' Remove a minimum 1 screen volume

c. Field Testing Equipment used:	Make	Model	Serial Number
	<u>YSI</u>	<u>556</u>	
	<u>Hanna</u>	<u>HI</u>	

Time (24hr)	Volume Removed (Liters)	Temp. (°C)	pH	Spec. Cond. (µS/cm)	DO (mg/L)	ORP (mV)	Turbidity (NTU)	Flow Rate (ml/min)	Drawdown (feet)	Color/Odor
<u>1000</u>										<u>Starch</u>
<u>1010</u>		<u>15.4</u>	<u>3.64</u>	<u>0.600</u>	<u>7.20</u>	<u>157.7</u>		<u>450</u>		
<u>1025</u>		<u>15.5</u>	<u>4.67</u>	<u>0.092</u>	<u>6.87</u>	<u>103.4</u>		<u>450</u>	<u>39.60</u>	
<u>1035</u>		<u>15.7</u>	<u>4.83</u>	<u>0.092</u>	<u>6.83</u>	<u>44.7</u>				
<u>1050</u>	<u>5 gal</u>			<u>40-50</u>	<u>over 100</u>					
<u>1110</u>				<u>back on</u>						
<u>1115</u>		<u>16.2</u>	<u>5.05</u>	<u>0.094</u>	<u>6.32</u>	<u>850</u>		<u>450</u>	<u>39.45</u>	

d. Acceptance criteria pass/fail Yes No N/A (continued on back)

- Has required volume been removed  Yes  No  N/A
- Has required turbidity been reached  Yes  No  N/A
- Have parameters stabilized  Yes  No  N/A

If no or N/A - Explain below.

purged 2 hours removed 9gals

### 3. SAMPLE COLLECTION: Method: bladder pump

Sample ID	Container Type	No. of Containers	Preservation	Analysis Req.	Time
<u>ED10801-GW-061014 10A</u>	<u>10A</u>	<u>3</u>	<u>HCl</u>	<u>VOCs</u>	<u>1010</u>
	<u>1-L water</u>	<u>2</u>	<u>-</u>	<u>14 Dioxane</u>	<u>1210</u>

Comments: sampled after 2 hrs parameters stable removed 9gal

Signature: Paul Kazeth Date: 6/10/14





RESOLUTION  
CONSULTANTS

Well ID: MW-10802

# Low Flow Ground Water Sample Collection Record

Client: Navy NWIRP Bethesda Date: 6/10/14 Time: Start 800 am/pm  
 Project No: 60256526 Finish 1200 am/pm  
 Site Location: Carl & Corona  
 Weather Conds: Cloudy, misting Collector(s): Pharaoh J. Christopher

### 1. WATER LEVEL DATA: (measured from Top of Casing)

a. Total Well Length 655 c. Length of Water Column 613.98 (a-b) Casing Diameter/Material 4" PVC  
 b. Water Table Depth 41.02 d. Calculated System Volume (see back) 400.8 gal

### 2. WELL PURGE DATA

a. Purge Method: Geotech bladder pump  
 b. Acceptance Criteria defined (see workplan)  
 - Temperature ± 3% - D.O. ± 10% (values >0.5 mg/L) Turbidity ± 10%  
 - pH ± 0.1 unit - ORP ± 10mV  
 - Sp. Cond. ± 3% - Drawdown < 0.3' Remove a minimum 1 screen volume

c. Field Testing Equipment used:

Make	Model	Serial Number
YSI	SS6 MPS	059240X
Hanna	HI 98703	065990X

Time (24hr)	Volume Removed (Liters)	Temp. (°C)	pH	Spec. Cond. (µS/cm)	DO (mg/L)	ORP (mV)	Turbidity (NTU)	Flow Rate (ml/min)	Drawdown (feet)	Color/Odor
905		18.85	6.62	0.114	7.06	153.2	4.89	250	40.98	Clear/None
910		16.97	4.33	0.094	6.21	272.1	4.76	500	40.99	Clear/None
915		15.58	3.36	0.080	5.73	347.0	4.66	500	40.92	Clear/None
920		15.42	3.38	0.079	5.21	348.9	4.79	500	40.90	Clear/None
925		15.59	3.97	0.078	4.76	323.0	3.16	400	40.87	Clear/None
930		15.57	4.14	0.077	4.58	302.6	2.54	400	40.85	Clear/None
1010		15.85	5.00	0.076	4.88	276.4	2.00	500	40.75	Clear/None

d. Acceptance criteria pass/fail (continued on back)

	Yes	No	N/A
Has required volume been removed	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Has required turbidity been reached	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Have parameters stabilized	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>

If no or N/A - Explain below.

### 3. SAMPLE COLLECTION: Method: Low flow

Sample ID	Container Type	No. of Containers	Preservation	Analysis Req.	Time
RE 10802 - GW 061014	ILAG	2	-	1,4 Dioxane	1145
RE 10802 - GW 061014	40ml VOA	3	HCl	VOA	1145

Comments \* Point issues @ 930

Signature [Signature] Date 6/10/14





RESOLUTION  
CONSULTANTS

Well ID: RA10301

# Low Flow Ground Water Sample Collection Record

Client: NWIRP Bathpage Date: 6/11/14 Time: Start 800 (am/pm) am  
 Project No: 60266526 Finish 1130 (am/pm) am  
 Site Location: Arceca & Martin  
 Weather Conds: Cloudy 65-70° Collector(s): P. Kersch

### 1. WATER LEVEL DATA: (measured from Top of Casing)

a. Total Well Length 645 c. Length of Water Column \_\_\_\_\_ (a-b) Casing Diameter/Material 4" PVC  
 b. Water Table Depth 40.52 d. Calculated System Volume (see back) 13 gal

### 2. WELL PURGE DATA

a. Purge Method: bladder pump  
 b. Acceptance Criteria defined (see workplan)  
 - Temperature ± 3% - D.O. ± 10% (values >0.5 mg/L) Turbidity ± 10%  
 - pH ± 0.1 unit - ORP ± 10mV  
 - Sp. Cond. ± 3% - Drawdown < 0.3' Remove a minimum 1 screen volume

c. Field Testing Equipment used:	Make	Model	Serial Number
	<u>YSI</u>	<u>556</u>	<u>1173912X</u>
	<u>Hanna</u>	<u>HI</u>	<u>4638908</u>

Time (24hr)	Volume Removed (Liters)	Temp. (°C)	pH	Spec. Cond. (µS/cm)	DO (mg/L)	ORP (mV)	Turbidity (NTU)	Flow Rate (ml/min)	Drawdown (feet)	Color/Odor
<u>855</u>										<u>01</u>
<u>900</u>		<u>16.8</u>	<u>499</u>	<u>121</u>	<u>7.81</u>	<u>320.1</u>				
<u>905</u>		<u>15.6</u>	<u>491</u>	<u>93</u>	<u>2.10</u>	<u>313</u>		<u>450</u>		
<u>910</u>			<u>493</u>	<u>93</u>	<u>2.27</u>	<u>300</u>				
<u>915</u>			<u>494</u>	<u>94</u>	<u>2.52</u>	<u>308</u>				
<u>920</u>		<u>15.0</u>	<u>495</u>	<u>95</u>	<u>2.49</u>	<u>301</u>			<u>40.15</u>	
<u>925</u>										

d. Acceptance criteria pass/fail (continued on back)

Has required volume been removed	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Has required turbidity been reached	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Have parameters stabilized	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>

If no or N/A - Explain below.

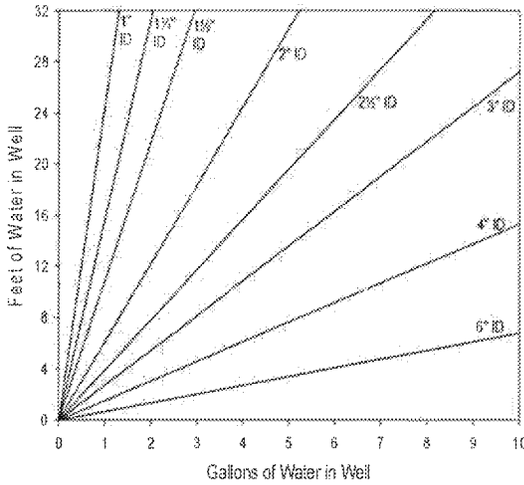
### 3. SAMPLE COLLECTION: Method: bladder pump

Sample ID	Container Type	No. of Containers	Preservation	Analysis Req.	Time
<u>RA10301-GW-061114</u>	<u>1/2 C vial</u>	<u>3</u>	<u>HCl</u>	<u>VOCs</u>	<u>1045</u>
<u>RA10301-GW-061114</u>	<u>1-L amber</u>	<u>2</u>	<u>-</u>	<u>1,4-Dioxane</u>	<u>1045</u>

Comments MS and MSD

Signature Paul Kersch Date 6/11/14

Purge Volume Calculation



Volume / Linear Ft. of Pipe		
ID (in)	Gallon	Liter
0.25	0.0025	0.0097
0.375	0.0057	0.0217
0.5	0.0102	0.0386
0.75	0.0229	0.0869
1	0.0408	0.1544
1.25	0.0637	0.2413
1.5	0.0918	0.3475
2	0.1632	0.6178
2.5	0.2550	0.9653
3	0.3672	1.3900
4	0.6528	2.4711
6	1.4688	5.5600

Well ID: 0N855 10301

(continued from front)

1 liter

Time (24 hr)	Volume Removed (Liters)	Temp (°C)	pH	Spec. Cond. (µS/cm)	DO (mg/L)	ORP (mV)	Turbidity (NTU)	Flow Rate (ml/min)	Drawdown (ft)	Color/Odor
930	500	15.0	4.72	103	4.15	322.0		450	39.95	
1040		15.0	4.70	103	4.27	322.8	0.48			
1145		15.0	4.72	103	4.21	322.6				
1230		15.0	4.73	102	4.26	322.3			39.65	
1355		15.0	4.74	102	4.25	321.8				
1400		15.1	4.74	102	4.26	321.2	0.30	450		
1505		15.1	4.76	101	4.32	319.8				
1610		15.1	4.76	101	4.30	317.6				
1715		15.2	4.77	101	4.28	316.5		450	39.45	
1820		15.3	4.81	101	4.29	311.1				
1925		15.2	4.81	100	4.27	305.7	1.16		39.45	
2030		15.3	4.84	100	4.29	302.6				
2135		15.3	4.83	100	4.31	297.5				
2240	1000	15.3	4.83	101	4.32	293.8	0.17	450		
1245										sample



Well ID: RE103 D3

RESOLUTION CONSULTANTS

# Low Flow Ground Water Sample Collection Record

Client: NWIRP Bethesda Date: 6/11/14 Time: Start 800 am/pm  
 Project No: 60266576 Finish 1400 am/pm  
 Site Location: Avoca & Martin  
 Weather Conds: Cloudy 65-70° Collector(s): Paul Kureth

### 1. WATER LEVEL DATA: (measured from Top of Casing)

a. Total Well Length 673 c. Length of Water Column \_\_\_\_\_ (a-b) Casing Diameter/Material 4" PVC  
 b. Water Table Depth 39.9 d. Calculated System Volume (see back) 13 gal <sup>Screen</sup>

### 2. WELL PURGE DATA

a. Purge Method: bladder pump  
 b. Acceptance Criteria defined (see workplan)  
 - Temperature ± 3% - D.O. ± 10% (values >0.5 mg/L) Turbidity ± 10%  
 - pH ± 0.1 unit - ORP ± 10mV  
 - Sp. Cond. ± 3% - Drawdown < 0.3' Remove a minimum 1 screen volume

c. Field Testing Equipment used:	Make	Model	Serial Number
	<u>YSI</u>	<u>556</u>	<u>1175912X</u>
	<u>Hanna</u>	<u>98703</u>	<u>1155810X</u>

Time (24hr)	Volume Removed (Liters)	Temp. (°C)	pH	Spec. Cond. (µS/cm)	DO (mg/L)	ORP (mV)	Turbidity (NTU)	Flow Rate (ml/min)	Drawdown (feet)	Color/Odor
1130								450	39.90	OK
1150		15.7	7.61	29	3.79	246.8	18.7			
1155		15.9	7.54	29	3.97	252.3				
1200		15.8	7.47	29	4.08	255.4				
1205		15.7	7.34	29	4.44	263.7			39.70	
1210	5 gal	15.7	7.24	30	4.51	274.0	3.07	450		
1215		15.6	7.24	20	4.55	275.4				

d. Acceptance criteria pass/fail (continued on back)

	Yes	No	N/A
Has required volume been removed	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Has required turbidity been reached	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Have parameters stabilized	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>

If no or N/A - Explain below.

### 3. SAMPLE COLLECTION: Method: bladder pump

Sample ID	Container Type	No. of Containers	Preservation	Analysis Req.	Time
<u>RE103D3-GW-061114</u>	<u>Vox vials</u>	<u>3</u>	<u>HCl</u>	<u>VOCs</u>	<u>1320</u>
<u>RE103D3-GW-061114</u>	<u>1-L amber</u>	<u>2</u>	<u>-</u>	<u>14 Dioxane</u>	<u>1320</u>

Comments \_\_\_\_\_

Signature Paul Kureth Date 06/11/14





Well ID: RE10502

RESOLUTION CONSULTANTS

# Low Flow Ground Water Sample Collection Record

Client: NWIRP Bethpage Date: 6/11/14 Time: Start 1400 am/pm  
 Project No: 60766576 Finish 1645 am/pm  
 Site Location: Roosevelt & Lincoln  
 Weather Conds: cloudy Collector(s): Paul Karath

### 1. WATER LEVEL DATA: (measured from Top of Casing)

a. Total Well Length 75592 c. Length of Water Column \_\_\_\_\_ (a-b) Casing Diameter/Material 4" PVC  
 b. Water Table Depth 37.54 d. Calculated System Volume (see back) 13 gal

### 2. WELL PURGE DATA

a. Purge Method: bladder pump

- b. Acceptance Criteria defined (see workplan)
- Temperature ± 3% - D.O. ± 10% (values >0.5 mg/L) Turbidity ± 10%
  - pH ± 0.1 unit - ORP ± 10mV
  - Sp. Cond. ± 3% - Drawdown < 0.3' Remove a minimum 1 screen volume

c. Field Testing Equipment used:

Make	Model	Serial Number
<u>XST</u>	<u>556</u>	<u>173912X</u>

Time (24hr)	Volume Removed (Liters)	Temp. (°C)	pH	Spec. Cond. (µS/cm)	DO (mg/L)	ORP (mV)	Turbidity (NTU)	Flow Rate (ml/min)	Drawdown (feet)	Color/Odor
<u>1445</u>										<u>oN</u>
<u>1500</u>		<u>15.9</u>	<u>5.00</u>	<u>56</u>	<u>4.74</u>	<u>217.3</u>		<u>400</u>		
<u>1510</u>		<u>15.7</u>	<u>4.97</u>	<u>56</u>	<u>4.62</u>	<u>213.3</u>				
<u>1510</u>		<u>15.7</u>	<u>4.97</u>	<u>56</u>	<u>4.55</u>	<u>212.5</u>		<u>500</u>		
<u>1515</u>		<u>15.7</u>	<u>4.97</u>	<u>56</u>	<u>3.86</u>	<u>206.6</u>	<u>1.76</u>		<u>37.40</u>	
<u>1520</u>		<u>16.1</u>	<u>4.93</u>	<u>57</u>	<u>3.34</u>	<u>204.6</u>				
<u>1525</u>		<u>16.3</u>	<u>4.98</u>	<u>59</u>	<u>3.54</u>	<u>200.1</u>				

d. Acceptance criteria pass/fail

	Yes	No	N/A
Has required volume been removed	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Has required turbidity been reached	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Have parameters stabilized	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>

If no or N/A - Explain below.

### 3. SAMPLE COLLECTION: Method: bladder pump

Sample ID	Container Type	No. of Containers	Preservation	Analysis Req.	Time
<u>RE10502-GW-061114</u>	<u>Marival</u>	<u>3</u>	<u>HCl</u>	<u>VOCs</u>	<u>1645</u>
<u>RE10502-GW-061114</u>	<u>1-L amber</u>	<u>2</u>	<u>-</u>	<u>14 Dioxane</u>	<u>1645</u>

Comments \_\_\_\_\_

Signature Paul Karath Date 6/11/14





RESOLUTION CONSULTANTS

Well ID: RE105D1

# Low Flow Ground Water Sample Collection Record

Client: NWIRP Bethpage Date: 6/11/14 Time: Start 1400 am/pm  
 Project No: 60260526 Finish 1700 am/pm  
 Site Location: Rosevelt Lincoln  
 Weather Conds: Cloudy Collector(s): G. Hicks

### 1. WATER LEVEL DATA: (measured from Top of Casing)

a. Total Well Length 538.9 c. Length of Water Column \_\_\_\_\_ (a-b) Casing Diameter/Material 4" PVC  
 b. Water Table Depth 36.50 d. Calculated System Volume (see back) 13 gal

### 2. WELL PURGE DATA

a. Purge Method: bladder pump  
 b. Acceptance Criteria defined (see workplan)  
 - Temperature ± 3% - D.O. ± 10% (values >0.5 mg/L) Turbidity ± 10%  
 - pH ± 0.1 unit - ORP ± 10mV  
 - Sp. Cond. ± 3% - Drawdown < 0.3' Remove a minimum 1 screen volume  
 c. Field Testing Equipment used: Make Model Serial Number  
QED Microprobe MPS0 MPS0-1112

Flow cell # 64432  
 YSI 556 mV: 54965

Time (24hr)	Volume Removed (Liters)	Temp. (°C)	pH	Spec. Cond. (µS/cm)	DO (mg/L)	ORP (mV)	Turbidity (NTU)	Flow Rate (ml/min)	Drawdown (feet)	Color/Odor
1435		18.24	5.37	176	5.49	-71.3	11.8	200	0	clear/none
1440		17.52	5.23	121	5.87	-78.9	10.9	300	"	"
1445		16.07	5.13	117	5.83	-65.1	2.66	"	"	"
1450		15.57	4.41	114	2.51	-96.9	0.97	"	"	"
1455		15.50	4.82	114	2.21	-100.1	0.82	"	"	"
1500		15.46	5.00	113	2.03	-128.2	0.86	"	"	"
1505		15.57	5.02	113	1.93	-128.0	1.02	350	"	"

d. Acceptance criteria pass/fail

	Yes	No	N/A
Has required volume been removed	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Has required turbidity been reached	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Have parameters stabilized	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>

If no or N/A - Explain below.

### 3. SAMPLE COLLECTION: Method: bladder pump

Sample ID	Container Type	No. of Containers	Preservation	Analysis Req.	Time
RE105D1-GW-061114	amb	2	-	14 Dioxane	1635
RE105D1-GW-061114	vac	3	HCl	VOC	1635
GW010-061114	amb	2	-	14 Dioxane	1645
GW010-061114	vac	3	HCl	VOC	1645

Comments \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

Signature: [Signature] Date: 6/11/14





RESOLUTION CONSULTANTS

Well ID: RE103D2

# Low Flow Ground Water Sample Collection Record

Client: NWIRP Bathpage Date: 6/11/14 Time: Start 9:00 am/pm  
 Project No: 60266566 Finish 11:00 am/pm  
 Site Location: Arco & Martin  
 Weather Conds: Cloudy 65-70 Collector(s): GH

### 1. WATER LEVEL DATA: (measured from Top of Casing)

a. Total Well Length 735 c. Length of Water Column \_\_\_\_\_ (a-b) Casing Diameter/Material 4" PVC  
 b. Water Table Depth 40.43 d. Calculated System Volume (see back) 13 gal

### 2. WELL PURGE DATA

a. Purge Method: Bladder pump  
 b. Acceptance Criteria defined (see workplan)  
 - Temperature ± 3% - D.O. ± 10% (values >0.5 mg/L) Turbidity ± 10%  
 - pH ± 0.1 unit - ORP ± 10mV  
 - Sp. Cond. ± 3% - Drawdown < 0.3' Remove a minimum 1 screen volume

c. Field Testing Equipment used:

Make	Model	Serial Number
YSI	556 MF	54965
Flow cell # 64432		

Time (24hr)	Volume Removed (Liters)	Temp. (°C)	pH	Spec. Cond. (µS/cm)	DO (mg/L)	ORP (mV)	Turbidity (NTU)	Flow Rate (ml/min)	Drawdown (feet)	Color/Odor
0840		15.56	3.92	35	10.72	86.5	8.61	200	0	clear/none
0845		15.52	3.99	37	7.15	63.7	12.51	"	0	"
0850		15.50	4.00	32	7.07	54.5	6.98	"	0	"
0855		15.45	4.02	43	7.12	41.5	1.99	"	0.07	"
0900		15.40	3.76	43	5.19	28.9	1.59	"	-	"
0905		15.39	4.17	41	5.04	41.4	14.2	"	0	"
0910		15.01	4.32	40	6.91	43.2	12.9	"	0	"

d. Acceptance criteria pass/fail

	Yes	No	N/A
Has required volume been removed	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Has required turbidity been reached	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Have parameters stabilized	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>

If no or N/A - Explain below.

### 3. SAMPLE COLLECTION: Method: bladder pump

Sample ID	Container Type	No. of Containers	Preservation	Analysis Req.	Time
RE103D2	amb	2	-	1,4 Dioxane	1040
RE103D2	VOA	3	HCl	VOC	1040

Comments \_\_\_\_\_

Signature: [Signature] Date: 6-11-14





RESOLUTION CONSULTANTS

Well ID: RE104D3

# Low Flow Ground Water Sample Collection Record

Client: NWIRP Bethesda Date: 6/12/14 Time: Start 800 am/pm  
 Project No: 60266576 Finish 1100 am/pm  
 Site Location: Hittop  
 Weather Conds: Cloudy 65-75° breeze Collector(s): Paul Kerech

### 1. WATER LEVEL DATA: (measured from Top of Casing)

a. Total Well Length 735 c. Length of Water Column \_\_\_\_\_ (a-b) Casing Diameter/Material 4" PVC  
 b. Water Table Depth 41.60 d. Calculated System Volume (see back) 13 gal

### 2. WELL PURGE DATA

a. Purge Method: bladder pump

b. Acceptance Criteria defined (see workplan)

- Temperature ± 3% - D.O. ± 10% (values >0.5 mg/L) Turbidity ± 10%
  - pH ± 0.1 unit - ORP ± 10mV
  - Sp. Cond. ± 3% - Drawdown < 0.3'
- Remove a minimum 1 screen volume

c. Field Testing Equipment used:	Make	Model	Serial Number
	<u>YSI</u>	<u>556</u>	
	<u>Hanna</u>	<u>HT</u>	

Time (24hr)	Volume Removed (Liters)	Temp. (°C)	pH	Spec. Cond. (µS/cm)	DO (mg/L)	ORP (mV)	Turbidity (NTU)	Flow Rate (ml/min)	Drawdown (feet)	Color/Odor
<u>810</u>									<u>41.60</u>	<u>ON</u>
<u>900</u>		<u>15.4</u>	<u>7.49</u>	<u>23</u>	<u>3.55</u>	<u>171.1</u>		<u>400</u>	<u>41.47</u>	
<u>920</u>		<u>15.3</u>	<u>7.56</u>	<u>21</u>	<u>4.54</u>	<u>174.0</u>		<u>400</u>	<u>41.85</u>	
<u>935</u>		<u>15.3</u>	<u>7.77</u>	<u>20</u>	<u>4.62</u>	<u>172.8</u>	<u>29.8</u>			
<u>930</u>		<u>15.4</u>	<u>7.81</u>	<u>20</u>	<u>4.61</u>	<u>173.7</u>				
<u>935</u>	<u>Spud</u>	<u>15.5</u>	<u>7.84</u>	<u>20</u>	<u>4.57</u>	<u>174.3</u>				
<u>940</u>		<u>15.6</u>	<u>7.88</u>	<u>20</u>	<u>4.50</u>	<u>175.0</u>	<u>21.5</u>			

d. Acceptance criteria pass/fail

	Yes	No	N/A
Has required volume been removed	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Has required turbidity been reached	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Have parameters stabilized	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>

If no or N/A - Explain below.

(continued on back)

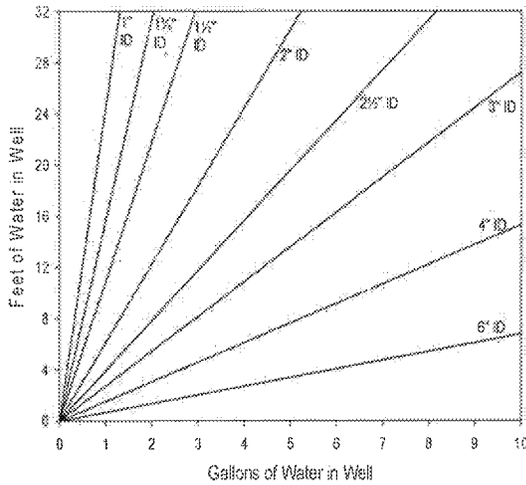
### 3. SAMPLE COLLECTION: Method: bladder pump

Sample ID	Container Type	No. of Containers	Preservation	Analysis Req.	Time
<u>RE10403-GW-061214</u>	<u>WV-Vial</u>	<u>3</u>	<u>HCl</u>	<u>VOL</u>	<u>1050</u>
<u>RE10409-GW-061214</u>	<u>VOL Vial</u>	<u>2</u>	<u>-</u>	<u>LA Trace</u>	<u>1050</u>

Comments \_\_\_\_\_

Signature Paul Kerech Date 6/12/14

Purge Volume Calculation



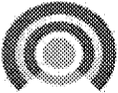
Volume / Linear Ft. of Pipe		
ID (in)	Gallon	Liter
0.25	0.0025	0.0097
0.375	0.0057	0.0217
0.5	0.0102	0.0386
0.75	0.0229	0.0869
1	0.0408	0.1544
1.25	0.0637	0.2413
1.5	0.0918	0.3475
2	0.1632	0.6178
2.5	0.2550	0.9653
3	0.3672	1.3900
4	0.6528	2.4711
6	1.4688	5.5600

Well ID: 944 DN

RE10403

(continued from front)

Time (24 hr)	Volume		Temp (°C)	pH	Spec. Cond. (µS/cm)	DO (mg/L)	ORP (mV)	Turbidity (NTU)	Flow Rate (ml/min)	Drawdown (ft)	Color/Odor
	Removed (Liters)										
945			15.5	8.94	20	4.49	175.6	11.9	100	41.3	
950			15.5	8.98	20	4.43	172.5				
955			15.5	8.96	20	4.41	175.2				
1020			15.6	4.00	20	4.36	176.2				
1025			15.5	4.01	20	4.35	175.6	14.4		41.05	
1010			15.5	4.02	20	4.30	176.0				
1015	10 gal		15.6	4.03	20	4.26	175.0				
1020			15.5	4.07	20	4.24	175.6			42.97	
1025			15.6	4.09	20	4.15	175.0	17.3			
1030			15.6	4.08	20	4.13	175.6	12.5		42.75	
1035			15.6	4.09	20	4.09	175.6	14.8			
1040	8 gal		15.9	4.10	20	4.06	175.5	15.0			
1050											sample



RESOLUTION CONSULTANTS

# Low Flow Ground Water Sample Collection Record

Well ID: RE 104D2

Client: NWIRP Bellpage Date: 6/12/14 Time: Start 800 am/pm  
 Project No: 60266526 Finish 1435 am/pm  
 Site Location: Hilltop  
 Weather Conds: Cloudy 65-75° drizzle Collector(s): Paul Karath

### 1. WATER LEVEL DATA: (measured from Top of Casing)

a. Total Well Length 735 c. Length of Water Column \_\_\_\_\_ (a-b) Casing Diameter/Material 4" PVC  
 b. Water Table Depth 41.12 d. Calculated System Volume (see back) 13 gal

### 2. WELL PURGE DATA

a. Purge Method: bladder pump

b. Acceptance Criteria defined (see workplan)

- Temperature ± 3% - D.O. ± 10% (values >0.5 mg/L) Turbidity ± 10%
- pH ± 0.1 unit - ORP ± 10mV
- Sp. Cond. ± 3% - Drawdown < 0.3' Remove a minimum 1 screen volume

c. Field Testing Equipment used:

Make	Model	Serial Number
<u>VST</u>	<u>556</u>	<u>59965</u>
<u>Hanna</u>	<u>HJ</u>	

Time (24hr)	Volume Removed (Liters)	Temp. (°C)	pH	Spec. Cond. (µS/cm)	DO (mg/L)	ORP (mV)	Turbidity (NTU)	Flow Rate (ml/min)	Drawdown (feet)	Color/Odor
<u>1245</u>										<u>19.1</u>
<u>1247</u>		<u>15.8</u>	<u>4.68</u>	<u>24</u>	<u>8.11</u>	<u>167.4</u>		<u>500</u>	<u>39.87</u>	
<u>1310</u>		<u>15.4</u>	<u>4.71</u>	<u>23</u>	<u>5.37</u>	<u>-7.9</u>				
<u>1320</u>	<u>5 gal</u>	<u>15.4</u>	<u>4.77</u>	<u>20</u>	<u>5.32</u>	<u>7.3</u>	<u>1.02</u>	<u>500</u>	<u>38.68</u>	
<u>1325</u>		<u>15.5</u>	<u>4.72</u>	<u>20</u>	<u>5.14</u>	<u>15.4</u>				
<u>1330</u>		<u>15.6</u>	<u>4.74</u>	<u>20</u>	<u>5.00</u>	<u>6.3</u>	<u>10.8</u>			
<u>1335</u>		<u>15.5</u>	<u>4.71</u>	<u>20</u>	<u>4.93</u>	<u>6.5</u>				

d. Acceptance criteria pass/fail (continued on back)

- |                                     | Yes                                 | No                                  | N/A                      |
|-------------------------------------|-------------------------------------|-------------------------------------|--------------------------|
| Has required volume been removed    | <input checked="" type="checkbox"/> | <input type="checkbox"/>            | <input type="checkbox"/> |
| Has required turbidity been reached | <input checked="" type="checkbox"/> | <input type="checkbox"/>            | <input type="checkbox"/> |
| Have parameters stabilized          | <input checked="" type="checkbox"/> | <input checked="" type="checkbox"/> | <input type="checkbox"/> |

If no or N/A - Explain below.

problem with ORP probe

### 3. SAMPLE COLLECTION: Method: bladder pump

Sample ID	Container Type	No. of Containers	Preservation	Analysis Req.	Time
<u>RE 104D2-GW-061214</u>	<u>100 vials</u>	<u>3</u>	<u>HCl</u>	<u>VOCs</u>	<u>1435</u>
<u>RE 104D2-GW-061214</u>	<u>1-L amber</u>	<u>2</u>	<u>-</u>	<u>1,4 Dioxane</u>	<u>1435</u>

Comments \_\_\_\_\_

Signature Paul Karath Date 6/12/14





RESOLUTION CONSULTANTS

Well ID: RE104D1

# Low Flow Ground Water Sample Collection Record

Client: NWIRP Belltop Date: 6/12/14 Time: Start 1050 am/pm  
 Project No: 60266526 Finish 1330 am/pm  
 Site Location: Hilltop Collector(s): G. Hicks  
 Weather Conds: Cloudy 65-75° drizzle

### 1. WATER LEVEL DATA: (measured from Top of Casing)

a. Total Well Length 375 c. Length of Water Column \_\_\_\_\_ (a-b) Casing Diameter/Material 4" PVC  
 b. Water Table Depth 35.78 d. Calculated System Volume (see back) 13 gal

### 2. WELL PURGE DATA

a. Purge Method: bladder pump  
 b. Acceptance Criteria defined (see workplan)  
 - Temperature ± 3% - D.O. ± 10% (values >0.5 mg/L) Turbidity ± 10%  
 - pH ± 0.1 unit - ORP ± 10mV  
 - Sp. Cond. ± 3% - Drawdown < 0.3' Remove a minimum 1 screen volume

c. Field Testing Equipment used:

Make	Model	Serial Number
YSI	560 MP	U52240A
H1 98703	Turbidimeter	U63890K
QED (22937) Well Wizard and QED MP50		

Time (24hr)	Volume Removed (Liters)	Temp. (°C)	pH	Spec. Cond. (µS/cm)	DO (mg/L)	ORP (mV)	Turbidity (NTU)	Flow Rate (ml/min)	Drawdown (feet)	Color/Odor
1050		16.00	6.78	69	2.65	185.0	1.11	350	0	clear / none
1055		15.98	5.57	69	8.28	192.1	0.89	400	0	"
1100		15.67	5.20	70	12.42	212.8	4.62	"	0	"
1105		15.66	4.95	66	6.28	313.8	11.12	"	0	"
1110		15.52	4.62	67	3.12	279.7	31.51	"	0	"
1115		15.49	4.61	66	4.87	262.5	6.20	"	0	"
1120	11	15.43	4.60	65	4.29	249.5	1.15	"	0	"

d. Acceptance criteria pass/fail

	Yes	No	N/A
Has required volume been removed	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Has required turbidity been reached	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Have parameters stabilized	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>

If no or N/A - Explain below.

### 3. SAMPLE COLLECTION: Method: bladder pump

Sample ID	Container Type	No. of Containers	Preservation	Analysis Req.	Time
RE104D1-GW-061214	VQA/amb	3/2	HCl/none	VOC/14 Dioxane	1750

Comments \_\_\_\_\_

Signature [Signature] Date 6-12-14





Well ID: BPOWS-2

# Low Flow Ground Water Sample Collection Record

Client: NWIRP Bethpage Date: 6/13/14 Time: Start 8:00 am/pm  
 Project No: 60266526 Finish 12:00 am/pm  
 Site Location: Killdare Crescent  
 Weather Conds: Foggy 70° Collector(s): GH

### 1. WATER LEVEL DATA: (measured from Top of Casing)

a. Total Well Length 585 c. Length of Water Column \_\_\_\_\_ (a-b) Casing Diameter/Material 4" PVC  
 b. Water Table Depth 21.22 d. Calculated System Volume (see back) 13 gal

### 2. WELL PURGE DATA

a. Purge Method: Bladder Pump

b. Acceptance Criteria defined (see workplan)

- Temperature ± 3% - D.O. ± 10% (values >0.5 mg/L) Turbidity ± 10%
- pH ± 0.1 unit - ORP ± 10mV
- Sp. Cond. ± 3% - Drawdown < 0.3' Remove a minimum 1 screen volume

c. Field Testing Equipment used:

Make	Model	Serial Number
<u>YSI 556 mps</u>	<u>U52240X</u>	<u>163896</u>
<u>Hanna</u>	<u>HT</u>	

Time (24hr)	Volume Removed (Liters)	Temp. (°C)	pH	Spec. Cond. (µS/cm)	DO (mg/L)	ORP (mV)	Turbidity (NTU)	Flow Rate (ml/min)	Drawdown (feet)	Color/Odor
0915		15.81	7.36	75	12.18	229.8	32.12	400	0	Clear/none
0920		16.90	4.87	74	4.81	234.6	12.40	"	"	"
0925	4	17.10	4.78	74	3.32	231.5	8.97	"	"	"
0930		17.05	4.42	67	3.11	235.7	8.01	"	"	"
0935		17.10	4.41	74	2.60	249.2	8.85	"	"	"
0940		17.15	4.41	79	1.51	248.5	7.12	"	"	"
0945		17.19	4.41	79	1.51	250.0	8.19	"	"	"

d. Acceptance criteria pass/fail

	Yes	No	N/A
Has required volume been removed	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Has required turbidity been reached	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Have parameters stabilized	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>

If no or N/A - Explain below.

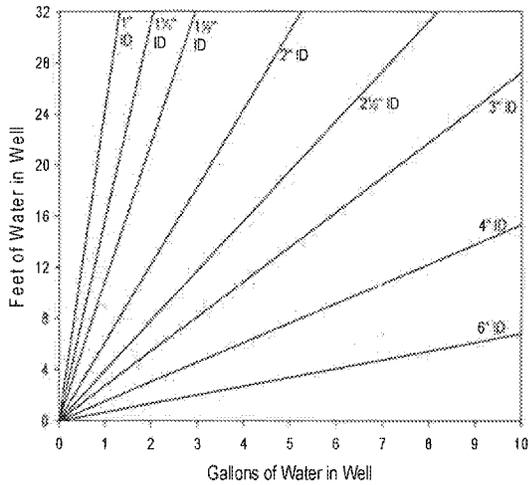
### 3. SAMPLE COLLECTION: Method: bladder pump

Sample ID	Container Type	No. of Containers	Preservation	Analysis Req.	Time
<u>BPOWS-2-GW-061314</u>	<u>VOA/amb</u>	<u>3/2</u>	<u>HCl/none</u>	<u>VOC/1,4 Dioxane</u>	<u>11:20</u>

Comments \_\_\_\_\_

Signature [Signature] Date 6-13-14

Purge Volume Calculation



Volume / Linear Ft. of Pipe		
ID (in)	Gallon	Liter
0.25	0.0025	0.0097
0.375	0.0057	0.0217
0.5	0.0102	0.0386
0.75	0.0229	0.0869
1	0.0408	0.1544
1.25	0.0637	0.2413
1.5	0.0918	0.3475
2	0.1632	0.6178
2.5	0.2550	0.9653
3	0.3672	1.3900
4	0.6528	2.4711
6	1.4688	5.5600

Well ID:

(continued from front)

09/15 Start

Time (24 hr)	Volume Removed (Liters)	Temp (°C)	pH	Spec. Cond. (µS/cm)	DO (mg/L)	ORP (mV)	Turbidity (NTU)	Flow Rate (ml/min)	Drawdown (ft)	Color/Odor
1000		15.98	4.47	72	1.02	240.2	24.6	400	0	clear/none
1015		16.72	4.51	71	0.98	189.7	68.3	400	"	"
1030	71	16.82	4.73	73	1.22	108.6	71,000	"	"	"
1045		16.00	5.86	105	5.93	50.1	>1,000	200	0	Cloudy/none
1010		17.98	5.71	88	2.79	78.3	71,000	"	"	"
1015		17.75	5.68	81	2.10	100.7	>1,000	"	"	"
1020		16.25	5.39	80	1.11	106.2	>1,000	"	"	"
1025		16.29	5.27	80	0.92	98.2	71,000	300	0	"
1030		16.31	5.10	79	0.67	84.3	>1,000	450	"	"
1035		16.11	5.25	84	0.55	69.1	71,000	450	"	"
1040		16.09	4.33	73	0.70	126.6	71,000	"	0.1	"
1045		16.15	4.20	74	0.66	135.9	52.7	"	0	"
1050		16.25	4.17	75	0.68	136.5	61.4	"	"	Clear/none
1055	42	16.26	4.12	76	0.67	138.9	33.3	"	"	"
1100		16.20	4.05	76	0.64	139.0	16.3	"	"	"
1105		16.20	4.05	76	0.63	125.1	12.2	"	"	"
1110		16.19	4.05	76	0.63	135.1	9.66	"	"	"
1115		16.19	4.08	76	0.62	131.8	8.97	"	"	"
1120	52	16.19	4.07	76	0.63	130.7	8.78	"	"	"



RESOLUTION CONSULTANTS

Well ID: BPOWS-1

# Low Flow Ground Water Sample Collection Record

Client: NOIRP Bethpage Date: 6/13/14 Time: Start 800 am/pm  
 Project No: 60266526 Finish 1400 am/pm  
 Site Location: Killdam Crescent  
 Weather Conds: Cloudy 70° Collector(s): Gordon Hicks

### 1. WATER LEVEL DATA: (measured from Top of Casing)

a. Total Well Length 515 c. Length of Water Column \_\_\_\_\_ (a-b) Casing Diameter/Material 4" PVC  
 b. Water Table Depth 20.92 d. Calculated System Volume (see back) 13 gal

### 2. WELL PURGE DATA

a. Purge Method: Bladder pump  
 b. Acceptance Criteria defined (see workplan)  
 - Temperature ± 3% - D.O. ± 10% (values >0.5 mg/L) Turbidity ± 10%  
 - pH ± 0.1 unit - ORP ± 10mV  
 - Sp. Cond. ± 3% - Drawdown < 0.3' Remove a minimum 1 screen volume

c. Field Testing Equipment used:

Make	Model	Serial Number
<u>Hanna</u>	<u>YSI 556 MPS</u>	<u>463890</u>
	<u>YSI 2240X</u>	

Time (24hr)	Volume Removed (Liters)	Temp. (°C)	pH	Spec. Cond. (µS/cm)	DO (mg/L)	ORP (mV)	Turbidity (NTU)	Flow Rate (ml/min)	Drawdown (feet)	Color/Odor
1140		18.99	5.61	101	12.56	212.1	65.12	350	0	Clear / none
1145		18.76	5.49	97	6.79	176.5	31.75	400	0	clear / none
1150		18.64	5.40	97	3.09	145.7	10.81	400	0	Clear / none
1155		18.20	4.78	96	1.24	139.0	2.32	"	"	"
1200	4	18.76	4.55	97	1.22	152.7	8.33	450	0	"
1205		18.70	4.41	98	1.23	175.1	7.93	"	"	"
1210		18.16	4.65	98	1.14	174.5	6.27	"	"	"

d. Acceptance criteria pass/fail (continued on back)

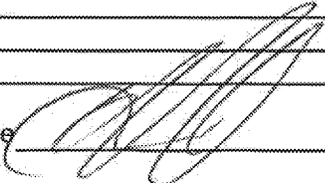
	Yes	No	N/A
Has required volume been removed	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Has required turbidity been reached	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Have parameters stabilized	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>

If no or N/A - Explain below.

### 3. SAMPLE COLLECTION: Method: bladder pump

Sample ID	Container Type	No. of Containers	Preservation	Analysis Req.	Time
<u>BPOWS-1-GW-061314</u>	<u>VOC/AMB</u>	<u>3/2</u>	<u>HCl/acet</u>	<u>VOC/1,4 dioxane</u>	<u>1330</u>

Comments \_\_\_\_\_

Signature:  Date: 6-13-14





RESOLUTION CONSULTANTS

Well ID: BPOW-3

# Low Flow Ground Water Sample Collection Record

Client: NWIRP Bathpage Date: 6/13/14 Time: Start 800 (am/pm) am  
 Project No: 60266526 Finish 1130 (am/pm) pm  
 Site Location: Killdeer Crescent  
 Weather Conds: Foggy 71° Collector(s): Paul Kareth

### 1. WATER LEVEL DATA: (measured from Top of Casing)

a. Total Well Length 665 c. Length of Water Column \_\_\_\_\_ (a-b) Casing Diameter/Material 4" PVC  
 b. Water Table Depth 21.34 d. Calculated System Volume (see back) 13 gal

### 2. WELL PURGE DATA

a. Purge Method: bladder pump  
 b. Acceptance Criteria defined (see workplan)  
 - Temperature ± 3% - D.O. ± 10% (values >0.5 mg/L) Turbidity ... ± 10%  
 - pH ± 0.1 unit - ORP ± 10mV  
 - Sp. Cond. ± 3% - Drawdown < 0.3' Remove a minimum 1 screen volume

c. Field Testing Equipment used:	Make	Model	Serial Number
	<u>YSI</u>	<u>556</u>	<u>74965</u>
	<u>Hanna</u>	<u>HI</u>	<u>11638902</u>

Time (24hr)	Volume Removed (Liters)	Temp. (°C)	pH	Spec. Cond. (µS/cm)	DO (mg/L)	ORP (mV)	Turbidity (NTU)	Flow Rate (ml/min)	Drawdown (feet)	Color/Odor
920										OK
930		16.0	2.67	33	1.64	196.7		550		
940		16.2	3.37	32	1.24	103.0	21000		21.25	
945		15.9	4.59	44	1.10	98.7				
950	Spot	15.8	4.08	30	0.96	132.9			21.83	
955										clean flow cell
100		16.1	4.50	32	1.72	140.9	268	550	21.95	

d. Acceptance criteria pass/fail

	Yes	No	N/A
Has required volume been removed	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Has required turbidity been reached	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
Have parameters stabilized	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>

If no or N/A - Explain below.

### 3. SAMPLE COLLECTION: Method: bladder pump

Sample ID	Container Type	No. of Containers	Preservation	Analysis Req.	Time
<u>BPOW-3-GW-061314</u>	<u>varials</u>	<u>3</u>	<u>HC1</u>	<u>VOC</u>	<u>1100</u>
<u>BPOW-3-GW-061314</u>	<u>varials</u>	<u>2</u>	<u>-</u>	<u>140 min</u>	<u>1100</u>

Comments pump kicked bottom up about 10:55, caused water to  
be so high up at 21.95  
rain at 9:45, pulled transducer during sampling

Signature Paul Kareth Date 6/13/14



**Appendix B**  
**Analytical Lab Sheets**

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4223-2  
**Client ID:** RE103D1-GW-061114  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4223  
**Lab File ID:** C7660.D

**Sample Date:** 11-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 14-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144769

**Analysis Date:** 14-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 17-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	J	0.54	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	J	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	J	0.25	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene		9.4	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113		18	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane		1.3	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene		4.2	ug/L	1	1	1.0	0.21	0.50
Chloroform	J	0.95	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	J	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	J	0.54	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	J	1000	ug/L	20	1	20	0.285	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	UM	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	J	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	J	0.68	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene		4.9	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	J	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

R 9/2/14

## Report of Analytical Results

Client: ENSAFE  
Lab ID: SH4223-2  
Client ID: RE103D1-GW-061114  
Project: Navy Clean WE15-03-06 NW  
SDG: SH4223  
Lab File ID: C7660.D

Sample Date: 11-JUN-14  
Received Date: 13-JUN-14  
Extract Date: 14-JUN-14  
Extracted By: REC  
Extraction Method: SW846 5030  
Lab Prep Batch: WG144769

Analysis Date: 14-JUN-14  
Analyst: REC  
Analysis Method: SW846 8260B  
Matrix: AQ  
% Solids: NA  
Report Date: 17-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	<del>U</del> <i>US</i>	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
<b>1,2-Dichloroethylene (Total)</b>		4.2	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		91.5	%					
Toluene-d8		93.1	%					
1,2-Dichloroethane-d4		113.	%					
Dibromofluoromethane		98.1	%					

*St 0/12/14*

### Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4223-1  
**Client ID:** RE103D2-GW-061114  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4223  
**Lab File ID:** C7659.D

**Sample Date:** 11-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 14-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144769

**Analysis Date:** 14-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 17-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U <i>UJ</i>	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
<b>1,1-Dichloroethene</b>	J	0.61	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
<b>Freon-113</b>		3.8	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
<b>1,1-Dichloroethane</b>	J	0.67	ug/L	1	1	1.0	0.21	0.50
<b>cis-1,2-Dichloroethene</b>		1.1	ug/L	1	1	1.0	0.21	0.50
<b>Chloroform</b>	J	0.96	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U <i>UJ</i>	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
<b>Trichloroethene</b>	U <i>J</i>	670	ug/L	1/10	1	1/10	0.28 2-8	0.70 5.0
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U <i>UJ</i>	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
<b>1,1,2-Trichloroethane</b>	J	0.37	ug/L	1	1	1.0	0.33	0.50
<b>Tetrachloroethene</b>	J	0.77	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U <i>UJ</i>	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*8/9/2/14*

### Report of Analytical Results

Client: ENSAFE  
Lab ID: SH4223-1  
Client ID: RE103D2-GW-061114  
Project: Navy Clean WE15-03-06 NW  
SDG: SH4223  
Lab File ID: C7659.D

Sample Date: 11-JUN-14  
Received Date: 13-JUN-14  
Extract Date: 14-JUN-14  
Extracted By: REC  
Extraction Method: SW846 5030  
Lab Prep Batch: WG144769

Analysis Date: 14-JUN-14  
Analyst: REC  
Analysis Method: SW846 8260B  
Matrix: AQ  
% Solids: NA  
Report Date: 17-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U <i>JS</i>	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
<b>1,2-Dichloroethylene (Total)</b>	J	1.1	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		93.2	%					
Toluene-d8		96.0	%					
1,2-Dichloroethane-d4		114.	%					
Dibromofluoromethane		99.2	%					

*Rec 6/17/14*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4223-5  
**Client ID:** RE103D3-GW-061114  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4223  
**Lab File ID:** C7663.D

**Sample Date:** 11-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 14-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144769

**Analysis Date:** 14-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 17-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	<del>U</del> U <sup>5</sup>	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
<b>1,1-Dichloroethene</b>	J	0.71	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
<b>Freon-113</b>		2.9	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
<b>1,1-Dichloroethane</b>	J	0.64	ug/L	1	1	1.0	0.21	0.50
<b>cis-1,2-Dichloroethene</b>	J	0.87	ug/L	1	1	1.0	0.21	0.50
<b>Chloroform</b>	J	0.94	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	<del>U</del> U <sup>5</sup>	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
<b>Carbon Tetrachloride</b>	J	0.30	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
<b>Trichloroethene</b>	<del>U</del> J	<del>5.0</del> 5.0	ug/L	<del>1</del> 10	1	<del>10</del> 10	<del>0.28</del> 2.8	<del>0.50</del> 5.0
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	<del>U</del> U <sup>5</sup>	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	<del>U</del> U <sup>5</sup>	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

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### Report of Analytical Results

Client: ENSAFE  
 Lab ID: SH4223-5  
 Client ID: RE103D3-GW-061114  
 Project: Navy Clean WE15-03-06 NW  
 SDG: SH4223  
 Lab File ID: C7663.D

Sample Date: 11-JUN-14  
 Received Date: 13-JUN-14  
 Extract Date: 14-JUN-14  
 Extracted By: REC  
 Extraction Method: SW846 5030  
 Lab Prep Batch: WG144769

Analysis Date: 14-JUN-14  
 Analyst: REC  
 Analysis Method: SW846 8260B  
 Matrix: AQ  
 % Solids: NA  
 Report Date: 17-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	<del>U</del> <i>US</i>	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
<b>1,2-Dichloroethylene (Total)</b>	J	0.87	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		95.4	%					
Toluene-d8		96.7	%					
1,2-Dichloroethane-d4		119.	%					
Dibromofluoromethane		105.	%					

*R 5/12/14*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4223-10  
**Client ID:** RE104D1-GW-061214  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4223  
**Lab File ID:** C7668.D

**Sample Date:** 12-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 14-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144769

**Analysis Date:** 14-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 17-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	J J	0.98	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	J UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
<b>1,1-Dichloroethene</b>		1.5	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
<b>Freon-113</b>		6.7	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
<b>1,1-Dichloroethane</b>	J	0.44	ug/L	1	1	1.0	0.21	0.50
<b>cis-1,2-Dichloroethene</b>		1.6	ug/L	1	1	1.0	0.21	0.50
<b>Chloroform</b>	J	0.37	ug/L	1	1	1.0	0.32	0.50
<b>1,1,1-Trichloroethane</b>	J	0.41	ug/L	1	1	1.0	0.20	0.50
2-Butanone	J UJ	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
<b>Trichloroethene</b>		160	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	J UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
<b>Tetrachloroethene</b>		2.4	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	J UJ	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*REC 6/12/14*

### Report of Analytical Results

Client: ENSAFE  
 Lab ID: SH4223-10  
 Client ID: RE104D1-GW-061214  
 Project: Navy Clean WE15-03-06 NW  
 SDG: SH4223  
 Lab File ID: C7668.D

Sample Date: 12-JUN-14  
 Received Date: 13-JUN-14  
 Extract Date: 14-JUN-14  
 Extracted By: REC  
 Extraction Method: SW846 5030  
 Lab Prep Batch: WG144769

Analysis Date: 14-JUN-14  
 Analyst: REC  
 Analysis Method: SW846 8260B  
 Matrix: AQ  
 % Solids: NA  
 Report Date: 17-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	<del>U</del> <i>WS</i>	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
<b>1,2-Dichloroethylene (Total)</b>	J	1.6	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		91.1	%					
Toluene-d8		92.7	%					
1,2-Dichloroethane-d4		113.	%					
Dibromofluoromethane		98.6	%					

*REC 6/12/14*

### Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4223-11  
**Client ID:** RE104D2-GW-061214  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4223  
**Lab File ID:** C7690.D

**Sample Date:** 12-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 16-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144799

**Analysis Date:** 16-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 17-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U UT	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U UT	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U J	0.84	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U UT	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U J	1.8	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U UT	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U UT	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U UT	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

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## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4223-11  
**Client ID:** RE104D2-GW-061214  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4223  
**Lab File ID:** C7690.D

**Sample Date:** 12-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 16-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144799

**Analysis Date:** 16-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 17-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	✓ UJ	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
<b>1,2-Dichloroethylene (Total)</b>	✓ UJ	0.84	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		92.4	%					
Toluene-d8		96.0	%					
1,2-Dichloroethane-d4		114.	%					
Dibromofluoromethane		99.2	%					

6/12/14

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4223-9  
**Client ID:** RE104D3-GW-061214  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4223  
**Lab File ID:** C7689.D

**Sample Date:** 12-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 16-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144799

**Analysis Date:** 16-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 17-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U <i>US</i>	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U <i>US</i>	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U <i>US</i>	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U <i>US</i>	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U <i>US</i>	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U <i>US</i>	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*US/12/14*

**Report of Analytical Results**

Client: ENSAFE  
Lab ID: SH4223-9  
Client ID: RE104D3-GW-061214  
Project: Navy Clean WE15-03-06 NW  
SDG: SH4223  
Lab File ID: C7689.D

Sample Date: 12-JUN-14  
Received Date: 13-JUN-14  
Extract Date: 16-JUN-14  
Extracted By: REC  
Extraction Method: SW846 5030  
Lab Prep Batch: WG144799

Analysis Date: 16-JUN-14  
Analyst: REC  
Analysis Method: SW846 8260B  
Matrix: AQ  
% Solids: NA  
Report Date: 17-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U <i>4.5</i>	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		90.9	%					
Toluene-d8		94.2	%					
1,2-Dichloroethane-d4		112.	%					
Dibromofluoromethane		97.1	%					

*8/9/12/14*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4223-6  
**Client ID:** RE105D1-GW-061114  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4223  
**Lab File ID:** C7664.D

**Sample Date:** 11-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 14-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144769

**Analysis Date:** 14-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 17-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	J J	1.4	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	J UJ	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
<b>1,1-Dichloroethene</b>		1.5	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
<b>Freon-113</b>		12	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
<b>1,1-Dichloroethane</b>	J	0.39	ug/L	1	1	1.0	0.21	0.50
<b>cis-1,2-Dichloroethene</b>		1.9	ug/L	1	1	1.0	0.21	0.50
<b>Chloroform</b>	J	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	J UJ	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
<b>Carbon Tetrachloride</b>	J	0.24	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
<b>Trichloroethene</b>		130	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	J UJ	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	J UJ	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

R. S. L. / 1/14

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4223-6  
**Client ID:** RE105D1-GW-061114  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4223  
**Lab File ID:** C7664.D

**Sample Date:** 11-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 14-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144769

**Analysis Date:** 14-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 17-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	<del>U</del> <i>UJ</i>	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
<b>1,2-Dichloroethylene (Total)</b>	J	1.9	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		92.4	%					
Toluene-d8		94.6	%					
1,2-Dichloroethane-d4		114.	%					
Dibromofluoromethane		99.6	%					

*Esther*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4223-7  
**Client ID:** RE105D2-GW-061114  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4223  
**Lab File ID:** C7665.D

**Sample Date:** 11-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 14-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144769

**Analysis Date:** 14-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 17-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	J J	0.58	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	J U J	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene		6.2	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113		32	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane		1.5	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene		3.1	ug/L	1	1	1.0	0.21	0.50
Chloroform		2.3	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	J	0.71	ug/L	1	1	1.0	0.20	0.50
2-Butanone	J U J	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride		4.7	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	J J	1200	ug/L	1/40	1	1040	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane		1.2	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	J	0.77	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	J U J	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

R9/12/14

## Report of Analytical Results

Client: ENSAFE  
Lab ID: SH4223-7  
Client ID: RE105D2-GW-061114  
Project: Navy Clean WE15-03-06 NW  
SDG: SH4223  
Lab File ID: C7665.D

Sample Date: 11-JUN-14  
Received Date: 13-JUN-14  
Extract Date: 14-JUN-14  
Extracted By: REC  
Extraction Method: SW846 5030  
Lab Prep Batch: WG144769

Analysis Date: 14-JUN-14  
Analyst: REC  
Analysis Method: SW846 8260B  
Matrix: AQ  
% Solids: NA  
Report Date: 17-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	<del>U</del> <i>UJ</i>	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
<b>1,2-Dichloroethylene (Total)</b>		3.1	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		91.5	%					
Toluene-d8		93.5	%					
1,2-Dichloroethane-d4		113.	%					
Dibromofluoromethane		98.8	%					

*ES/2/14*

### Report of Analytical Results

Client: ENSAFE  
 Lab ID: SH4223-3  
 Client ID: RE108D1-GW-061014  
 Project: Navy Clean WE15-03-06 NW  
 SDG: SH4223  
 Lab File ID: C7661.D

Sample Date: 10-JUN-14  
 Received Date: 13-JUN-14  
 Extract Date: 14-JUN-14  
 Extracted By: REC  
 Extraction Method: SW846 5030  
 Lab Prep Batch: WG144769

Analysis Date: 14-JUN-14  
 Analyst: REC  
 Analysis Method: SW846 8260B  
 Matrix: AQ  
 % Solids: NA  
 Report Date: 17-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U <i>UJ</i>	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	J	0.58	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U <i>UJ</i>	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene		82	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U <i>UJ</i>	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	J	0.59	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U <i>UJ</i>	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*REC 6/12/14*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4223-3  
**Client ID:** RE108D1-GW-061014  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4223  
**Lab File ID:** C7661.D

**Sample Date:** 10-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 14-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144769

**Analysis Date:** 14-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 17-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U <i>us</i>	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		91.9	%					
Toluene-d8		95.4	%					
1,2-Dichloroethane-d4		113.	%					
Dibromofluoromethane		98.8	%					

*Polizky*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4223-4  
**Client ID:** RE108D2-GW-061014  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4223  
**Lab File ID:** C7662.D

**Sample Date:** 10-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 14-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144769

**Analysis Date:** 14-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 17-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	J S	0.27	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	J S	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
<b>1,1-Dichloroethene</b>		8.2	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
<b>Freon-113</b>		8.7	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
<b>1,1-Dichloroethane</b>		5.8	ug/L	1	1	1.0	0.21	0.50
<b>cis-1,2-Dichloroethene</b>		9.9	ug/L	1	1	1.0	0.21	0.50
<b>Chloroform</b>		4.2	ug/L	1	1	1.0	0.32	0.50
<b>1,1,1-Trichloroethane</b>		1.3	ug/L	1	1	1.0	0.20	0.50
2-Butanone	J UT	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
<b>Carbon Tetrachloride</b>		2.0	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
<b>Trichloroethene</b>	J S	1800	3400 ug/L	1 40	1	10 40	0.28 11	0.50 20
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	J UT	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
<b>1,1,2-Trichloroethane</b>		1.8	ug/L	1	1	1.0	0.33	0.50
<b>Tetrachloroethene</b>		1.6	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	J UT	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4223-4  
**Client ID:** RE108D2-GW-061014  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4223  
**Lab File ID:** C7662.D

**Sample Date:** 10-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 14-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144769

**Analysis Date:** 14-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 17-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U <i>UJ</i>	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
<b>1,2-Dichloroethylene (Total)</b>		9.9	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		89.0	%					
Toluene-d8		91.3	%					
1,2-Dichloroethane-d4		112.	%					
Dibromofluoromethane		97.5	%					

*R 9/12/14*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4223-8  
**Client ID:** GWDUP-061114  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4223  
**Lab File ID:** C7666.D

**Sample Date:** 11-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 14-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144769

**Analysis Date:** 14-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 17-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	✓ J	1.4	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	✓ U J	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
<b>1,1-Dichloroethene</b>		1.7	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
<b>Freon-113</b>		13	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	I	6.8	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
<b>1,1-Dichloroethane</b>	J	0.40	ug/L	1	1	1.0	0.21	0.50
<b>cis-1,2-Dichloroethene</b>		1.9	ug/L	1	1	1.0	0.21	0.50
<b>Chloroform</b>	J	0.47	ug/L	1	1	1.0	0.32	0.50
<b>1,1,1-Trichloroethane</b>	J	0.62	ug/L	1	1	1.0	0.20	0.50
2-Butanone	✓ U J	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
<b>Carbon Tetrachloride</b>	J	0.27	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
<b>Trichloroethene</b>		140	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	✓ U J	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	✓ U J	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

R 9/2/14

## Report of Analytical Results

Client: ENSAFE  
Lab ID: SH4223-8  
Client ID: GWDUP-061114  
Project: Navy Clean WE15-03-06 NW  
SDG: SH4223  
Lab File ID: C7666.D

Sample Date: 11-JUN-14  
Received Date: 13-JUN-14  
Extract Date: 14-JUN-14  
Extracted By: REC  
Extraction Method: SW846 5030  
Lab Prep Batch: WG144769

Analysis Date: 14-JUN-14  
Analyst: REC  
Analysis Method: SW846 8260B  
Matrix: AQ  
% Solids: NA  
Report Date: 17-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	H UJ	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
<b>1,2-Dichloroethylene (Total)</b>	J	1.9	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		88.8	%					
Toluene-d8		91.5	%					
1,2-Dichloroethane-d4		114.	%					
Dibromofluoromethane		98.9	%					

*REC/12/14*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4223-12  
**Client ID:** RE-TB-061214  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4223  
**Lab File ID:** C7654.D

**Sample Date:** 12-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 14-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144769

**Analysis Date:** 14-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 17-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	U <i>US</i>	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
<b>Carbon Disulfide</b>	U <i>US</i>	<del>0.28</del> 1.0	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	U <i>US</i>	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	U <i>US</i>	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U <i>US</i>	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*RF 9/12/14*

### Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4223-12  
**Client ID:** RE-TB-061214  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4223  
**Lab File ID:** C7654.D

**Sample Date:** 12-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 14-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144769

**Analysis Date:** 14-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 17-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	<del>U</del> <i>U5</i>	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		95.2	%					
Toluene-d8		95.8	%					
1,2-Dichloroethane-d4		112.	%					
Dibromofluoromethane		99.1	%					

*6/12/14*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4223-2DL  
**Client ID:** RE103D1-GW-061114  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4223  
**Lab File ID:** G1817.D

**Sample Date:** 11-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 16-JUN-14  
**Extracted By:** AC  
**Extraction Method:** SW846 3520  
**Lab Prep Batch:** WG144790

**Analysis Date:** 20-JUN-14  
**Analyst:** JCG  
**Analysis Method:** SW846 M8270D  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
1,4-Dioxane	MM	20	ug/L	3	.25	0.77	0.26	0.56
1,4-Dioxane-D8		144.	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4223-1  
**Client ID:** RE103D2-GW-061114  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4223  
**Lab File ID:** G1797.D

**Sample Date:** 11-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 16-JUN-14  
**Extracted By:** AC  
**Extraction Method:** SW846 3520  
**Lab Prep Batch:** WG144790

**Analysis Date:** 19-JUN-14  
**Analyst:** JCG  
**Analysis Method:** SW846 M8270D  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
1,4-Dioxane		1.0	ug/L	1	.25	0.24	0.080	0.17
1,4-Dioxane-D8		120.	%					

### Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4223-5  
**Client ID:** RE103D3-GW-061114  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4223  
**Lab File ID:** G1803.D

**Sample Date:** 11-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 16-JUN-14  
**Extracted By:** AC  
**Extraction Method:** SW846 3520  
**Lab Prep Batch:** WG144790

**Analysis Date:** 19-JUN-14  
**Analyst:** JCG  
**Analysis Method:** SW846 M8270D  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
1,4-Dioxane		0.92	ug/L	1	.25	0.24	0.081	0.17
1,4-Dioxane-D8		121.	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4223-10DL  
**Client ID:** RE104D1-GW-061214  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4223  
**Lab File ID:** G1813.D

**Sample Date:** 12-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 16-JUN-14  
**Extracted By:** AC  
**Extraction Method:** SW846 3520  
**Lab Prep Batch:** WG144790

**Analysis Date:** 20-JUN-14  
**Analyst:** JCG  
**Analysis Method:** SW846 M8270D  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
1,4-Dioxane		12	ug/L	2	.25	0.47	0.16	0.34
1,4-Dioxane-D8		147.	%					

### Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4223-11  
**Client ID:** RE104D2-GW-061214  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4223  
**Lab File ID:** G1810.D

**Sample Date:** 12-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 16-JUN-14  
**Extracted By:** AC  
**Extraction Method:** SW846 3520  
**Lab Prep Batch:** WG144790

**Analysis Date:** 20-JUN-14  
**Analyst:** JCG  
**Analysis Method:** SW846 M8270D  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
1,4-Dioxane	U	0.17	ug/L	1	.25	0.24	0.081	0.17
1,4-Dioxane-D8		119.	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4223-9  
**Client ID:** RE104D3-GW-061214  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4223  
**Lab File ID:** G1807.D

**Sample Date:** 12-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 16-JUN-14  
**Extracted By:** AC  
**Extraction Method:** SW846 3520  
**Lab Prep Batch:** WG144790

**Analysis Date:** 19-JUN-14  
**Analyst:** JCG  
**Analysis Method:** SW846 M8270D  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
1,4-Dioxane	U	0.17	ug/L	1	.25	0.24	0.080	0.17
1,4-Dioxane-D8		107.	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4223-6DL  
**Client ID:** RE105D1-GW-061114  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4223  
**Lab File ID:** G1811.D

**Sample Date:** 11-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 16-JUN-14  
**Extracted By:** AC  
**Extraction Method:** SW846 3520  
**Lab Prep Batch:** WG144790

**Analysis Date:** 20-JUN-14  
**Analyst:** JCG  
**Analysis Method:** SW846 M8270D  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
1,4-Dioxane		15	ug/L	2	.25	0.50	0.17	0.36
1,4-Dioxane-D8		139.	%					

### Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4223-7  
**Client ID:** RE105D2-GW-061114  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4223  
**Lab File ID:** G1805.D

**Sample Date:** 11-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 16-JUN-14  
**Extracted By:** AC  
**Extraction Method:** SW846 3520  
**Lab Prep Batch:** WG144790

**Analysis Date:** 19-JUN-14  
**Analyst:** JCG  
**Analysis Method:** SW846 M8270D  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
1,4-Dioxane		6.2	ug/L	1	.25	0.24	0.080	0.17
1,4-Dioxane-D8		128.	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4223-3  
**Client ID:** RE108D1-GW-061014  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4223  
**Lab File ID:** G1801.D

**Sample Date:** 10-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 16-JUN-14  
**Extracted By:** AC  
**Extraction Method:** SW846 3520  
**Lab Prep Batch:** WG144790

**Analysis Date:** 19-JUN-14  
**Analyst:** JCG  
**Analysis Method:** SW846 M8270D  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
1,4-Dioxane		7.1	ug/L	1	.25	0.24	0.080	0.17
1,4-Dioxane-D8		111.	%					

### Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4223-4  
**Client ID:** RE108D2-GW-061014  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4223  
**Lab File ID:** G1802.D

**Sample Date:** 10-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 16-JUN-14  
**Extracted By:** AC  
**Extraction Method:** SW846 3520  
**Lab Prep Batch:** WG144790

**Analysis Date:** 19-JUN-14  
**Analyst:** JCG  
**Analysis Method:** SW846 M8270D  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
1,4-Dioxane		6.9	ug/L	1	.25	0.24	0.083	0.18
1,4-Dioxane-D8		107.	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4223-8DL  
**Client ID:** GWDUP-061114  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4223  
**Lab File ID:** G1812.D

**Sample Date:** 11-JUN-14  
**Received Date:** 13-JUN-14  
**Extract Date:** 16-JUN-14  
**Extracted By:** AC  
**Extraction Method:** SW846 3520  
**Lab Prep Batch:** WG144790

**Analysis Date:** 20-JUN-14  
**Analyst:** JCG  
**Analysis Method:** SW846 M8270D  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
1,4-Dioxane		18	ug/L	2	.25	0.52	0.18	0.37
1,4-Dioxane-D8		146.	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4293-3  
**Client ID:** BPOWS-1-GW-061314  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4293  
**Lab File ID:** C7734.D

**Sample Date:** 13-JUN-14  
**Received Date:** 17-JUN-14  
**Extract Date:** 18-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144955

**Analysis Date:** 18-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 19-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	<del>U</del> <i>US</i>	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	<del>U</del> <i>US</i>	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	<del>U</del> <i>US</i>	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	<del>U</del> <i>US</i>	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*Res/12/14*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4293-3  
**Client ID:** BPOWS-1-GW-061314  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4293  
**Lab File ID:** C7734.D

**Sample Date:** 13-JUN-14  
**Received Date:** 17-JUN-14  
**Extract Date:** 18-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144955

**Analysis Date:** 18-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 19-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		92.4	%					
Toluene-d8		96.8	%					
1,2-Dichloroethane-d4		115.	%					
Dibromofluoromethane		101.	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4293-2  
**Client ID:** BPOWS-2-GW-061314  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4293  
**Lab File ID:** C7733.D

**Sample Date:** 13-JUN-14  
**Received Date:** 17-JUN-14  
**Extract Date:** 18-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144955

**Analysis Date:** 18-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 19-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	H US	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	H US	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	H US	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	H US	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

*REC*

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4293-2  
**Client ID:** BPOWS-2-GW-061314  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4293  
**Lab File ID:** C7733.D

**Sample Date:** 13-JUN-14  
**Received Date:** 17-JUN-14  
**Extract Date:** 18-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144955

**Analysis Date:** 18-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 19-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		92.7	%					
Toluene-d8		96.7	%					
1,2-Dichloroethane-d4		114.	%					
Dibromofluoromethane		100.	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4293-1  
**Client ID:** BPOWS-3-GW-061314  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4293  
**Lab File ID:** C7732.D

**Sample Date:** 13-JUN-14  
**Received Date:** 17-JUN-14  
**Extract Date:** 18-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144955

**Analysis Date:** 18-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 19-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	H US	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	H US	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
Carbon Disulfide	U	0.50	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	H US	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	H US	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

R 9/17/14

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4293-1  
**Client ID:** BPOWS-3-GW-061314  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4293  
**Lab File ID:** C7732.D

**Sample Date:** 13-JUN-14  
**Received Date:** 17-JUN-14  
**Extract Date:** 18-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144955

**Analysis Date:** 18-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 19-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		93.2	%					
Toluene-d8		96.7	%					
1,2-Dichloroethane-d4		117.	%					
Dibromofluoromethane		102.	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4293-4  
**Client ID:** BPOWS-TB-061314  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4293  
**Lab File ID:** C7724.D

**Sample Date:** 13-JUN-14  
**Received Date:** 17-JUN-14  
**Extract Date:** 18-JUN-14  
**Extracted By:** REC  
**Extraction Method:** SW846 5030  
**Lab Prep Batch:** WG144955

**Analysis Date:** 18-JUN-14  
**Analyst:** REC  
**Analysis Method:** SW846 8260B  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 19-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Dichlorodifluoromethane	H US	1.0	ug/L	1	2	2.0	0.24	1.0
Chloromethane	U	1.0	ug/L	1	2	2.0	0.36	1.0
Vinyl Chloride	U	1.0	ug/L	1	2	2.0	0.25	1.0
Bromomethane	U	1.0	ug/L	1	2	2.0	0.49	1.0
Chloroethane	H US	1.0	ug/L	1	2	2.0	0.55	1.0
Trichlorofluoromethane	U	1.0	ug/L	1	2	2.0	0.24	1.0
1,1-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.35	0.50
<b>Carbon Disulfide</b>	J	0.35	ug/L	1	1	1.0	0.25	0.50
Freon-113	U	0.50	ug/L	1	1	1.0	0.31	0.50
Methylene Chloride	U	2.5	ug/L	1	5	5.0	1.1	2.5
Acetone	U	2.5	ug/L	1	5	5.0	2.2	2.5
trans-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.25	0.50
Methyl tert-butyl Ether	U	0.50	ug/L	1	1	1.0	0.36	0.50
1,1-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.21	0.50
cis-1,2-Dichloroethene	U	0.50	ug/L	1	1	1.0	0.21	0.50
Chloroform	U	0.50	ug/L	1	1	1.0	0.32	0.50
1,1,1-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
2-Butanone	H US	2.5	ug/L	1	5	5.0	1.3	2.5
Cyclohexane	U	0.50	ug/L	1	1	1.0	0.31	0.50
Carbon Tetrachloride	U	0.50	ug/L	1	1	1.0	0.22	0.50
Benzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,2-Dichloroethane	U	0.50	ug/L	1	1	1.0	0.20	0.50
Trichloroethene	U	0.50	ug/L	1	1	1.0	0.28	0.50
1,2-Dichloropropane	U	0.50	ug/L	1	1	1.0	0.25	0.50
Bromodichloromethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
cis-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.19	0.50
Toluene	U	0.50	ug/L	1	1	1.0	0.27	0.50
4-Methyl-2-Pentanone	H US	2.5	ug/L	1	5	5.0	1.3	2.5
trans-1,3-Dichloropropene	U	0.50	ug/L	1	1	1.0	0.20	0.50
1,1,2-Trichloroethane	U	0.50	ug/L	1	1	1.0	0.33	0.50
Tetrachloroethene	U	0.50	ug/L	1	1	1.0	0.40	0.50
Dibromochloromethane	U	0.50	ug/L	1	1	1.0	0.30	0.50
2-Hexanone	U	2.5	ug/L	1	5	5.0	1.7	2.5
Chlorobenzene	U	0.50	ug/L	1	1	1.0	0.22	0.50
Ethylbenzene	U	0.50	ug/L	1	1	1.0	0.21	0.50

## Report of Analytical Results

Client: ENSAFE  
Lab ID: SH4293-4  
Client ID: BPOWS-TB-061314  
Project: Navy Clean WE15-03-06 NW  
SDG: SH4293  
Lab File ID: C7724.D

Sample Date: 13-JUN-14  
Received Date: 17-JUN-14  
Extract Date: 18-JUN-14  
Extracted By: REC  
Extraction Method: SW846 5030  
Lab Prep Batch: WG144955

Analysis Date: 18-JUN-14  
Analyst: REC  
Analysis Method: SW846 8260B  
Matrix: AQ  
% Solids: NA  
Report Date: 19-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
Xylenes (total)	U	1.5	ug/L	1	3	3.0	0.25	1.5
Styrene	U	0.50	ug/L	1	1	1.0	0.23	0.50
Bromoform	U	0.50	ug/L	1	1	1.0	0.23	0.50
Isopropylbenzene	U	0.50	ug/L	1	1	1.0	0.23	0.50
1,1,2,2-Tetrachloroethane	U	0.50	ug/L	1	1	1.0	0.38	0.50
1,3-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.26	0.50
1,4-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.24	0.50
1,2-Dichlorobenzene	U	0.50	ug/L	1	1	1.0	0.15	0.50
1,2,4-Trichlorobenzene	U	0.50	ug/L	1	1	1.0	0.37	0.50
Methyl Acetate	U	0.75	ug/L	1	1	1.0	0.53	0.75
Methylcyclohexane	U	0.50	ug/L	1	1	1.0	0.30	0.50
o-Xylene	U	0.50	ug/L	1	1	1.0	0.25	0.50
M+P-Xylenes	U	1.0	ug/L	1	2	2.0	0.59	1.0
1,2-Dichloroethylene (Total)	U	1.0	ug/L	1	2	2.0	0.21	1.0
1,2-Dibromoethane	U	0.50	ug/L	1	1	1.0	0.22	0.50
1,2-Dibromo-3-Chloropropane	U	0.75	ug/L	1	1	1.0	0.50	0.75
P-Bromofluorobenzene		93.7	%					
Toluene-d8		95.7	%					
1,2-Dichloroethane-d4		114.	%					
Dibromofluoromethane		101.	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4293-1  
**Client ID:** BPOWS-3-GW-061314  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4293  
**Lab File ID:** G1822.D

**Sample Date:** 13-JUN-14  
**Received Date:** 17-JUN-14  
**Extract Date:** 19-JUN-14  
**Extracted By:** JMS  
**Extraction Method:** SW846 3520  
**Lab Prep Batch:** WG145030

**Analysis Date:** 20-JUN-14  
**Analyst:** JCG  
**Analysis Method:** SW846 M8270D  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
1,4-Dioxane		0.55	ug/L	1	.25	0.24	0.080	0.17
1,4-Dioxane-D8		112.	%					

## Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4293-2  
**Client ID:** BPOWS-2-GW-061314  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4293  
**Lab File ID:** G1823.D

**Sample Date:** 13-JUN-14  
**Received Date:** 17-JUN-14  
**Extract Date:** 19-JUN-14  
**Extracted By:** JMS  
**Extraction Method:** SW846 3520  
**Lab Prep Batch:** WG145030

**Analysis Date:** 20-JUN-14  
**Analyst:** JCG  
**Analysis Method:** SW846 M8270D  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
1,4-Dioxane	U	0.17	ug/L	1	.25	0.24	0.080	0.17
1,4-Dioxane-D8		116.	%					

### Report of Analytical Results

**Client:** ENSAFE  
**Lab ID:** SH4293-3  
**Client ID:** BPOWS-1-GW-061314  
**Project:** Navy Clean WE15-03-06 NW  
**SDG:** SH4293  
**Lab File ID:** G1824.D

**Sample Date:** 13-JUN-14  
**Received Date:** 17-JUN-14  
**Extract Date:** 19-JUN-14  
**Extracted By:** JMS  
**Extraction Method:** SW846 3520  
**Lab Prep Batch:** WG145030

**Analysis Date:** 20-JUN-14  
**Analyst:** JCG  
**Analysis Method:** SW846 M8270D  
**Matrix:** AQ  
**% Solids:** NA  
**Report Date:** 23-JUN-14

Compound	Qualifier	Result	Units	Dilution	LOQ	ADJ LOQ	ADJ MDL	ADJ LOD
1,4-Dioxane	U	0.17	ug/L	1	.25	0.24	0.080	0.17
1,4-Dioxane-D8		138.	%					

**Appendix C**  
**Data Validation**

## Data Validation Report

Project:	Regional Groundwater Investigation - NWIRP Bethpage	
Laboratory:	Katahdin Analytical Services, Inc.	
Service Request:	SH4223	
Analyses/Method:	EPA SW-846 Method 8260B for VOCs (GC/MS)	
Validation Level:	Limited	
RESCON Project Number:	60266526.SA.DV	
Prepared by:	Sheena Blair/RESCON	Completed on: 08/13/2014
Reviewed by:	Lori Herberich/RESCON	File Name: SH4223_8260B

### SUMMARY

The samples listed below were collected by Resolution Consultants (RESCON) from the Regional Groundwater Investigation - NWIRP Bethpage site on June 10, 11, and 12, 2014.

Sample ID	Matrix/Sample Type
GWDUP-061114	Field Duplicate of RE105D1-GW-061114
RE103D1-GW-061114	Ground water
RE103D2-GW-061114	Ground water
RE103D3-GW-061114	Ground water
RE104D1-GW-061214	Ground water
RE104D2-GW-061214	Ground water
RE104D3-GW-061214	Ground water
RE105D1-GW-061114	Ground water
RE105D2-GW-061114	Ground water
RE108D1-GW-061014	Ground water
RE108D2-GW-061014	Ground water
RE-TB-061214	Trip Blank

Data validation activities were conducted with reference to *Test Methods for Evaluating Solid Waste, Physical/Chemical Methods SW846, specifically SW-846 Method 8260B, Volatile Organic Compounds by Gas Chromatography/Mass Spectrometry* (USEPA, 1996), *USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review* (June 2008), and *Quality Systems Manual (QSM) for Environmental Laboratories, Version 4.2* (DoD, October 2010). In the absence of method-specific information, laboratory quality control (QC) limits, project-specific requirements and/or professional judgment were used as appropriate.

### REVIEW ELEMENTS

The data were evaluated based on the following review elements (where applicable to the method):

- ✓ Data completeness (chain-of-custody (COC))/sample integrity

- ✓ Holding times and sample preservation
- ✓ GC/MS performance checks
- X Initial calibration/continuing calibration verification
- X Laboratory blanks/trip blanks/equipment blanks
- ✓ Surrogate spike recoveries
- ✓ Matrix spike (MS) results
- ✓ Laboratory control sample (LCS) results
- ✓ Field duplicate results
- ✓ Internal standard results
- ✓ Sample results/reporting issues

The symbol (✓) indicates that no validation qualifiers were applied based on this parameter. NA indicates that the parameter was not included as part of this data set or was not applicable to this validation and therefore not reviewed. The symbol (X) indicates that a QC nonconformance resulted in the qualification of data. Any QC nonconformance that resulted in the qualification of data is discussed below. In addition, nonconformances or other issues that were noted during validation, but did not result in qualification of data, may be discussed for informational purposes only.

The data appear valid as reported and may be used for decision making purposes. Selected data points were estimated, negated, and/or rejected due to nonconformances of certain QC criteria (see discussion below). Qualified sample results are presented in Table 1.

## RESULTS

### Data Completeness/Sample Integrity

The data package was reviewed and found to meet acceptance criteria for completeness:

- The COCs were reviewed for completeness of information relevant to the samples and requested analyses, and for signatures indicating transfer of sample custody.
- The laboratory sample login sheet(s) were reviewed for issues potentially affecting sample integrity, including the condition of sample containers upon receipt at the laboratory.
- Completeness of analyses was verified by comparing the reported results to the COC requests.

Due to limitations in the reporting system, the laboratory truncated the ID for Trip Blank in the report to "RE-TB-061214. The submitted EDD file reflects the full sample ID.

### Holding Times/Sample Preservation

Sample preservation and preparation/analysis holding times were reviewed for conformance with the QC acceptance criteria.

The QC acceptance criteria were met.

### GC/MS Performance Checks

The data were reviewed to ensure that the 4-bromofluorobenzene (BFB) tuning was performed at the correct frequency and that the method acceptance criteria were met.

### Initial Calibration/Continuing Calibration Verification

Calibration data were reviewed for conformance with the QC acceptance criteria to ensure that:

- the initial calibration (ICAL) percent relative standard deviation (%RSD), correlation coefficient (r)/coefficient of determination ( $r^2$ ), and/or response factor method acceptance criteria were met;
- the initial calibration verification (ICV) percent recovery (%R) criteria were met; and
- the continuing calibration verification standard (CCV) method percent difference or percent drift (%Ds), %Rs, and/or RF acceptance criteria were met; and/or
- the retention time method acceptance criteria were met.

Data qualification to the analytes associated with the specific ICAL and/or CCV was as follows:

#### **ICAL Linearity Nonconformances:**

Nonconformance	Actions	
	Detected Results	Nondetected Results
%RSD > 15% and quantitation based on mean RF	J	UJ
r or $r^2$ < 0.99 and quantitation based on linear	J*	UJ*
* No guidance in NFG, thus AECOM professional judgment was used		

#### **ICV Recovery Nonconformances:**

Nonconformance	Actions	
	Detected Compounds	Nondetected Compounds
%R > 120%	J	No qualification
20% < %R < 80%	J	UJ
%R < 20% (see note)	J	R*

Notes: Based on NFG 2008 VOC guidance, professional judgment is used to reject (R) non-detects in all associated samples for any analyte with < 20% recovery. Also, professional judgment is used to estimate (UJ) rather the reject (R) sample results previously negated (U) on the basis of blank contamination.

#### **CCV Linearity Nonconformances:**

Nonconformance	Actions	
	Detected Results	Nondetected Results
%D > 20%	J	UJ
%Drift > 20%	J*	UJ*
* No guidance in NFG, thus professional judgment was used		

Qualified sample results are shown in Table 1. Nonconformances are summarized in Attachment A in Tables A-1a, A-1b, and A-1c.

### Laboratory Blanks/Trip Blanks/Equipment Blanks

Laboratory method blanks, equipment rinsate and trip blanks were evaluated as to whether there were contaminants detected above the detection limit (DL).

Data validation qualifications for individual samples are based on the maximum contaminant concentration detected in all associated blanks.

Method, equipment rinsate and trip blank results were reviewed for conformance with the QC acceptance criteria. Detected results in blanks are not discussed in this data validation report if the associated results were nondetect or if qualification of sample results was not required.

Sample results were qualified as follows:

Blank type	Blank result	Sample result	Action for samples
Method, Storage, Field, Trip, or Instrument*	Detects	Not detected	No qualification
	≤ LOQ	< LOQ	Report sample LOQ value with a U
		> LOQ and ≤ 2x LOQ	Report the sample result with a U**
		≥ 2x the LOQ	No qualifications
	> LOQ	< LOQ	Report sample LOQ value with a U
		≥ LOQ and < blank contamination	Report the sample result with a U or reject the sample result as unusable R
		≥ LOQ and ≥ blank contamination	If the result is ≤ 2x blank result, report the sample result U.** If the result is > 2x blank result, no qualification is required.**
TIC detected	Detects	If the result is ≤ 2x blank result, report the sample result U.** If the result is > 2x blank result, no qualification is required.**	
* Qualifications based on instrument blank results affect only the sample analyzed immediately after the sample that has target compounds that exceed the calibration range or non-target compounds that exceed 100 g/L.			
**Based on RESCON professional judgment.			

LOQ - Limit of Quantitation

Nonconformances are summarized in Attachment A in Table A-2. Qualified sample results are shown in Table 1.

### Surrogate Spike Recoveries

The surrogate recoveries (%Rs) were reviewed for conformance with the QC acceptance criteria.

All QC acceptance criteria were met.

### MS Results

The MS %Rs were reviewed for conformance with the QC acceptance criteria.

The MS %R for bromodichloromethane was slightly above the upper QC limit. However, the sample was nondetect, and the result was accepted without qualification.

### LCS Results

The LCS %Rs were reviewed for conformance with the QC acceptance criteria.

All QC acceptance criteria were met.

#### **Field Duplicate Results**

Field duplicate relative percent differences (RPDs) were reviewed for conformance with the QC criterion of  $\leq 30\%$  for aqueous matrices. This criterion applies if both results were greater than five times the Limit of Quantitation (LOQ).

All QC acceptance criteria were met.

#### **Internal Standard Results**

The internal standard (IS) recoveries were reviewed for conformance with the QC acceptance criteria.

All QC acceptance criteria were met.

#### **Sample Results/Reporting Issues**

Compounds that were not detected in the sample are reported as undetected (U) at the Limit of Detection (LOD).

Compounds detected at concentrations less than the LOQ but greater than the DL were qualified by the laboratory as estimated (J). This "J" qualifier was retained during data validation.

Any sample that was analyzed at a dilution due to high concentrations of target or non-target compounds or matrix interferences was checked to ensure that the results and/or sample specific LODs and LOQs were adjusted accordingly by the laboratory.

#### **QUALIFICATION ACTIONS**

Sample results qualified as a result of validation actions are summarized in Table 1. All actions are described above.

#### **ATTACHMENTS**

Attachment A: Nonconformance Summary Tables

Attachment B: Qualifier Codes and Explanations

Attachment C: Reason Codes and Explanations

Table 1 - Data Validation Summary of Qualified Data

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
GWDUP-061114	WG	2-BUTANONE		2.5	ug/L	UJ	c
GWDUP-061114	WG	2-HEXANONE		2.5	ug/L	UJ	c
GWDUP-061114	WG	4-METHYL-2-PENTANONE		2.5	ug/L	UJ	c
GWDUP-061114	WG	ACETONE	6.8	2.5	ug/L	J	c
GWDUP-061114	WG	BROMOFORM		0.50	ug/L	UJ	c
GWDUP-061114	WG	CHLOROETHANE		1.0	ug/L	UJ	c
GWDUP-061114	WG	DICHLORODIFLUOROMETHANE	1.4	1.0	ug/L	J	c
RE103D1-GW-061114	WG	2-BUTANONE		2.5	ug/L	UJ	c
RE103D1-GW-061114	WG	2-HEXANONE		2.5	ug/L	UJ	c
RE103D1-GW-061114	WG	4-METHYL-2-PENTANONE		2.5	ug/L	UJ	c
RE103D1-GW-061114	WG	BROMOFORM		0.50	ug/L	UJ	c
RE103D1-GW-061114	WG	CHLOROETHANE		1.0	ug/L	UJ	c
RE103D1-GW-061114	WG	DICHLORODIFLUOROMETHANE	0.54	1.0	ug/L	J	c
RE103D1-GW-061114	WG	TRICHLOROETHENE	1200	10	ug/L	J	c
RE103D2-GW-061114	WG	2-BUTANONE		2.5	ug/L	UJ	c
RE103D2-GW-061114	WG	2-HEXANONE		2.5	ug/L	UJ	c
RE103D2-GW-061114	WG	4-METHYL-2-PENTANONE		2.5	ug/L	UJ	c
RE103D2-GW-061114	WG	BROMOFORM		0.50	ug/L	UJ	c
RE103D2-GW-061114	WG	CHLOROETHANE		1.0	ug/L	UJ	c
RE103D2-GW-061114	WG	TRICHLOROETHENE	670	5.0	ug/L	J	c
RE103D3-GW-061114	WG	2-BUTANONE		2.5	ug/L	UJ	c
RE103D3-GW-061114	WG	2-HEXANONE		2.5	ug/L	UJ	c
RE103D3-GW-061114	WG	4-METHYL-2-PENTANONE		2.5	ug/L	UJ	c
RE103D3-GW-061114	WG	BROMOFORM		0.50	ug/L	UJ	c
RE103D3-GW-061114	WG	CHLOROETHANE		1.0	ug/L	UJ	c
RE103D3-GW-061114	WG	TRICHLOROETHENE	510	5.0	ug/L	J	c
RE104D1-GW-061214	WG	2-BUTANONE		2.5	ug/L	UJ	c
RE104D1-GW-061214	WG	2-HEXANONE		2.5	ug/L	UJ	c
RE104D1-GW-061214	WG	4-METHYL-2-PENTANONE		2.5	ug/L	UJ	c
RE104D1-GW-061214	WG	BROMOFORM		0.50	ug/L	UJ	c
RE104D1-GW-061214	WG	CHLOROETHANE		1.0	ug/L	UJ	c
RE104D1-GW-061214	WG	DICHLORODIFLUOROMETHANE	0.98	1.0	ug/L	J	c
RE104D2-GW-061214	WG	1,2-DICHLOROETHENE, TOTAL	0.84	1.0	ug/L	J	c
RE104D2-GW-061214	WG	4-METHYL-2-PENTANONE		2.5	ug/L	UJ	c
RE104D2-GW-061214	WG	BROMOFORM		0.50	ug/L	U	c
RE104D2-GW-061214	WG	CARBON TETRACHLORIDE		0.50	ug/L	UJ	c
RE104D2-GW-061214	WG	CHLOROETHANE		1.0	ug/L	U	c
RE104D2-GW-061214	WG	CIS-1,2-DICHLOROETHENE	0.84	0.50	ug/L	J	c
RE104D2-GW-061214	WG	CIS-1,3-DICHLOROPROPENE		0.50	ug/L	UJ	c

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
RE104D2-GW-061214	WG	DICHLORODIFLUOROMETHANE		1.0	ug/L	UJ	c
RE104D2-GW-061214	WG	TRANS-1,3-DICHLOROPROPENE		0.50	ug/L	UJ	c
RE104D2-GW-061214	WG	TRICHLOROETHENE	1.8	0.50	ug/L	J	c
RE104D3-GW-061214	WG	4-METHYL-2-PENTANONE		2.5	ug/L	UJ	c
RE104D3-GW-061214	WG	BROMOFORM		0.50	ug/L	U	c
RE104D3-GW-061214	WG	CARBON TETRACHLORIDE		0.50	ug/L	UJ	c
RE104D3-GW-061214	WG	CHLOROETHANE		1.0	ug/L	U	c
RE104D3-GW-061214	WG	CIS-1,3-DICHLOROPROPENE		0.50	ug/L	UJ	c
RE104D3-GW-061214	WG	DICHLORODIFLUOROMETHANE		1.0	ug/L	UJ	c
RE104D3-GW-061214	WG	TRANS-1,3-DICHLOROPROPENE		0.50	ug/L	UJ	c
RE105D1-GW-061114	WG	2-BUTANONE		2.5	ug/L	UJ	c
RE105D1-GW-061114	WG	2-HEXANONE		2.5	ug/L	UJ	c
RE105D1-GW-061114	WG	4-METHYL-2-PENTANONE		2.5	ug/L	UJ	c
RE105D1-GW-061114	WG	BROMOFORM		0.50	ug/L	UJ	c
RE105D1-GW-061114	WG	CHLOROETHANE		1.0	ug/L	UJ	c
RE105D1-GW-061114	WG	DICHLORODIFLUOROMETHANE	1.4	1.0	ug/L	J	c
RE105D2-GW-061114	WG	2-BUTANONE		2.5	ug/L	UJ	c
RE105D2-GW-061114	WG	2-HEXANONE		2.5	ug/L	UJ	c
RE105D2-GW-061114	WG	4-METHYL-2-PENTANONE		2.5	ug/L	UJ	c
RE105D2-GW-061114	WG	BROMOFORM		0.50	ug/L	UJ	c
RE105D2-GW-061114	WG	CHLOROETHANE		1.0	ug/L	UJ	c
RE105D2-GW-061114	WG	DICHLORODIFLUOROMETHANE	0.58	1.0	ug/L	J	c
RE105D2-GW-061114	WG	TRICHLOROETHENE	1500	20	ug/L	J	c
RE108D1-GW-061014	WG	2-BUTANONE		2.5	ug/L	UJ	c
RE108D1-GW-061014	WG	2-HEXANONE		2.5	ug/L	UJ	c
RE108D1-GW-061014	WG	4-METHYL-2-PENTANONE		2.5	ug/L	UJ	c
RE108D1-GW-061014	WG	BROMOFORM		0.50	ug/L	UJ	c
RE108D1-GW-061014	WG	CHLOROETHANE		1.0	ug/L	UJ	c
RE108D2-GW-061014	WG	2-BUTANONE		2.5	ug/L	UJ	c
RE108D2-GW-061014	WG	2-HEXANONE		2.5	ug/L	UJ	c
RE108D2-GW-061014	WG	4-METHYL-2-PENTANONE		2.5	ug/L	UJ	c
RE108D2-GW-061014	WG	BROMOFORM		0.50	ug/L	UJ	c
RE108D2-GW-061014	WG	CHLOROETHANE		1.0	ug/L	UJ	c
RE108D2-GW-061014	WG	DICHLORODIFLUOROMETHANE	0.27	1.0	ug/L	J	c
RE108D2-GW-061014	WG	TRICHLOROETHENE	3400	20	ug/L	J	c
RE-TB-061214	WQ	2-BUTANONE		2.5	ug/L	UJ	c
RE-TB-061214	WQ	2-HEXANONE		2.5	ug/L	UJ	c
RE-TB-061214	WQ	4-METHYL-2-PENTANONE		2.5	ug/L	UJ	c
RE-TB-061214	WQ	BROMOFORM		0.50	ug/L	UJ	c
RE-TB-061214	WQ	CARBON DISULFIDE		1.0*	ug/L	U	bl

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
RE-TB-061214	WQ	CHLOROETHANE		1.0	ug/L	UJ	c
*Carbon disulfide -LOQ							

## Attachment A

## Nonconformance Summary Tables

Table A-1a - Initial Calibration Verification

ICAL	Compound	% RSD	Limit
GCMS-C	CHLOROETHANE	36.2	<15%
	BROMOFORM	15.5	<15%
ICV	Compound	% R	Limit
WG141592	DICHLOROFLUOROMETHANE	142.1	80-120%
	ACETONE	120.4	80-120%

Associated samples: All samples in the SDG

Table A-1b - Continuing Calibration Verification

CCV	Compound	% D	Limit
WG144769	CHLOROETHANE	22.1	<20%
	2-BUTANONE	20.5	<20%
	4-METHYL-2-PENTANONE	37.1	<20%
	2-HEXANONE	24.5	<20%

Associated samples: GWDUP-061114, RE-TB-061214, RE103D1-GW-061114, RE103D2-GW-061114, RE103D3-GW-061114, RE104D1-GW-061214, RE105D1-GW-061114, RE105D2-GW-061114, RE108D1-GW-061014, RE108D2-GW-061014

Table A-1c - Continuing Calibration Verification

CCV	Compound	% D	Limit
WG144799	DICHLOROFLUOROMETHANE	74.6	<20%
	CARBON TETRACHLORIDE	22.1	<20%
	CIS-1,3-DICHLOROPROPENE	21.2	<20%
	4-METHYL-2-PENTANONE	28.3	<20%
	TRANS-1,3-DICHLOROPROPENE	20.6	<20%

Associated samples: RE104D2-GW-061214, RE104D3-GW-061214

Table A-2 - Lab Blanks

Blank ID	Compound	Result	LOD	Units	Associated Samples
WG144769-2	CARBON DISULFIDE	0.30	0.50	ug/L	RE-TB-061214

**Attachment B****Qualifier Codes and Explanations**

<b>Qualifier</b>	<b>Explanation</b>
J	The analyte was positively identified; the associated numerical value is the approximate concentration of the analyte in the sample.
UJ	The analyte was not detected above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte in the sample.
U	The analyte was analyzed for, but was not detected above the reported sample quantitation limit.
R	The sample results are rejected due to serious deficiencies in the ability to analyze the sample and meet quality control criteria. The presence or absence of the analyte cannot be verified.

## Attachment C

## Reason Codes and Explanations

Reason Code	Explanation
be	Equipment blank contamination
bf	Field blank contamination
bf	Laboratory blank contamination
c	Calibration issue
co	Analyte carryover
d	Reporting limit raised due to chromatographic interference
fd	Field duplicate RPDs
h	Holding times
i	Internal standard areas
k	Estimated Maximum Possible Concentration (EMPC)
l	LCS or OPR recoveries
lc	Labeled compound recovery
ld	Laboratory duplicate RPDs
lp	Laboratory control sample/laboratory control sample duplicate RPDs
m	Matrix spike recovery
md	Matrix spike/matrix spike duplicate RPDs
nb	Negative laboratory blank contamination
p	Chemical preservation issue
r	Dual column RPD
q	Quantitation issue
s	Surrogate recovery
su	Ion suppression
t	Temperature preservation issue
x	Percent solids
y	Serial dilution results
z	ICS results

## Data Validation Report

Project:	Regional Groundwater Investigation - NWIRP Bethpage	
Laboratory:	Katahdin Analytical Services, Inc.	
Service Request:	SH4223	
Analyses/Method:	EPA SW-846 Method 8270C-SIM for SVOCs (GC/MS)	
Validation Level:	Limited	
RESCON Project Number:	60266526.SA.DV	
Prepared by:	Sheena Blair/RESCON	Completed on: 08/13/2014
Reviewed by:	Lori Herberich/RESCON	File Name: SH4223 SW7270D

### SUMMARY

The samples listed below were collected by Resolution Consultants (RESCON) from the Regional Groundwater Investigation - NWIRP Bethpage site on June 10, 11, and 12, 2014.

Sample ID	Matrix/Sample Type
GWDUP-061114	Field Duplicate of RE105D2-GW-061114
RE103D1-GW-061114	Ground water
RE103D2-GW-061114	Ground water
RE103D3-GW-061114	Ground water
RE104D1-GW-061214	Ground water
RE104D2-GW-061214	Ground water
RE104D3-GW-061214	Ground water
RE105D1-GW-061114	Ground water
RE105D2-GW-061114	Ground water
RE108D1-GW-061014	Ground water
RE108D2-GW-061014	Ground water

Data validation activities were conducted with reference to *Test Methods for Evaluating Solid Waste, Physical/Chemical Methods SW846, specifically SW-846 Method 8260B, Volatile Organic Compounds by Gas Chromatography/Mass Spectrometry* (USEPA, 1996), *USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review* (June 2008), and *Quality Systems Manual (QSM) for Environmental Laboratories, Version 4.2* (DoD, October 2010). In the absence of method-specific information, laboratory quality control (QC) limits, project-specific requirements and/or professional judgment were used as appropriate.

### REVIEW ELEMENTS

The data were evaluated based on the following review elements (where applicable to the method):

- ✓ Data completeness (chain-of-custody (COC)/sample integrity)
- ✓ Holding times and sample preservation

- ✓ Instrument tuning
- ✓ Initial calibration/continuing calibration verification
- ✓ Laboratory blanks/equipment blanks
- ✓ Surrogate spike recoveries
- ✓ Matrix spike (MS) and/or matrix spike duplicate (MSD) results
- ✓ Laboratory control sample (LCS)/laboratory control sample duplicate (LCSD) results
- ✓ Field duplicate results
- ✓ Internal standard results
- ✓ Sample results/reporting issues

The symbol (✓) indicates that no results were qualified based on this parameter. NA indicates that the parameter was not included as part of this data set or was not applicable to this validation and therefore not reviewed. The symbol (X) indicates that a quality control (QC) nonconformance resulted in the qualification of data. Any QC nonconformance that resulted in the qualification of data is discussed below. In addition, nonconformances or other issues that were noted during validation, but did not result in qualification of data, may be discussed for informational purposes only.

The data appear valid as reported and may be used for decision making purposes. There were no results qualified on the basis of this data review.

## RESULTS

### Data Completeness

The data package was reviewed and found to meet acceptance criteria for completeness:

- The COCs were reviewed for completeness of information relevant to the samples and requested analyses, and for signatures indicating transfer of sample custody.
- The laboratory sample login sheet(s) were reviewed for issues potentially affecting sample integrity, including the condition of sample containers upon receipt at the laboratory.
- Completeness of analyses was verified by comparing the reported results to the COC requests.

### Holding Times/Sample Preservation

Sample preservation and preparation/analysis holding times were reviewed for conformance with the QC acceptance criteria.

The QC acceptance criteria were met.

### GC/MS Performance Checks

The data were reviewed to ensure that the decafluorotriphenylphosphine (DFTPP) tuning was performed at the correct frequency and that the method acceptance criteria were met. All samples were analyzed within 12 hours of the DFTPP tunes.

### **Initial Calibration/Continuing Calibration Verification**

Calibration data were reviewed for conformance with the QC acceptance criteria to ensure that:

- the initial calibration (ICAL) percent relative standard deviation (%RSD), correlation coefficient (r)/coefficient of determination ( $r^2$ ), and/or response factor method acceptance criteria were met;
- the initial calibration verification (ICV) percent recovery (%R) criteria were met; and
- the continuing calibration verification standard (CCV) method percent difference or percent drift (%Ds), %Rs, and/or RF acceptance criteria were met; and/or
- the retention time method acceptance criteria were met.

The QC acceptance criteria were met.

### **Laboratory Blanks/Equipment Blanks**

Laboratory method blanks and equipment rinsate blanks were evaluated as to whether there were contaminants detected above the detection limit (DL).

Data validation qualifications for individual samples are based on the maximum contaminant concentration detected in all associated blanks.

Detected results in blanks are not discussed in this data validation report if the associated results were nondetect or if qualification of sample results was not required.

### **Surrogate Spike Recoveries**

The surrogate recoveries (%Rs) were reviewed for conformance with the QC acceptance criteria.

All QC acceptance criteria were met.

### **MS/MSD Results**

The MS/MSD %Rs and/or relative percent differences (RPDs) were reviewed for conformance with the QC acceptance criteria.

The concentrations of 1,4-dioxane of in the native sample and the MS/MSD were outside the calibration curve and the %Rs could not be calculated. The RPD between the MS and MSD met the QC limits. No validation action was taken on this basis.

### **LCS/LCSD Results**

The LCS/LCSD %Rs and/or RPDs were reviewed for conformance with the QC acceptance criteria.

All QC acceptance criteria were met.

### **Field Duplicate Results**

Field duplicate RPDs were reviewed for conformance with the AECOM QC criteria of  $\leq 30\%$  for aqueous matrices. This criteria applies if both results were greater than five times the quantitation limit (QL).

All QC acceptance criteria were met and/or qualification of the data was not required.

#### **Internal Standard Results**

The internal standard (IS) recoveries were reviewed for conformance with the QC acceptance criteria.

All QC acceptance criteria were met.

#### **Sample Results/Reporting Issues**

All compounds detected at concentrations less than the quantitation limit (LOQ) but greater than the DL were qualified by the laboratory as estimated (J). This "J" qualifier was retained during data validation.

Any sample that was analyzed at a dilution due to high concentrations of target or non-targets was checked to ensure that the results and/or sample specific QLs were adjusted accordingly by the laboratory.

#### **QUALIFICATION ACTIONS**

No sample results were qualified as a result of this data review.

#### **ATTACHMENTS**

Attachment A: Nonconformance Summary Tables

Attachment B: Qualifier Codes and Explanations

**Attachment A**

**Nonconformance Summary Tables**

No nonconformances were identified in this data set.

**Attachment B****Qualifier Codes and Explanations**

<b>Qualifier</b>	<b>Explanation</b>
J	The analyte was positively identified; the associated numerical value is the approximate concentration of the analyte in the sample.
UJ	The analyte was not detected above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte in the sample.
U	The analyte was analyzed for, but was not detected above the reported sample quantitation limit.
R	The sample results are rejected due to serious deficiencies in the ability to analyze the sample and meet quality control criteria. The presence or absence of the analyte cannot be verified.



## Data Validation Report

Project:	Regional Groundwater Investigation - NWIRP Bethpage	
Laboratory:	Katahdin Analytical Services, Scarborough, Maine	
Service Request:	SH4293	
Analyses/Method:	EPA SW-846 Method 8260B for VOCs (GC/MS) and EPA SW-846 Method 8270C-SIM for SVOCs (GC/MS)	
Validation Level:	Limited	
RESCON Project Number:	60266526.SA.DV	
Prepared by:	Sheena Blair/RESCON	Completed on: 9/12/2014
Reviewed by:	Lori Herberich/RESCON	File Name: SH4293_8260B and 8270D_SIM

### SUMMARY

The samples listed below were collected by Resolution Consultants (RESCON) from the Regional Groundwater Investigation - NWIRP Bethpage site on June 13, 2014.

Sample ID	Matrix/Sample Type
BPOW5-1-GW-061314	Groundwater
BPOW5-2-GW-061314	Groundwater
BPOW5-3-GW-061314	Groundwater
BPOW5-TB-061314	Trip Blank

Data validation activities were conducted with reference to *Test Methods for Evaluating Solid Waste, Physical/Chemical Methods SW846, specifically SW-846 Method 8260B, Volatile Organic Compounds by Gas Chromatography/Mass Spectrometry* (USEPA, 1996), *Test Methods for Evaluating Solid Waste, Physical/Chemical Methods SW846, specifically SW-846 Method 8270D-SIM, Semivolatile Organic Compounds by Gas Chromatography/Mass Spectrometry* (USEPA, 1996); *USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review* (June 2008), and *Quality Systems Manual (QSM) for Environmental Laboratories, Version 4.2* (DoD, October 2010). In the absence of method-specific information, laboratory quality control (QC) limits, project-specific requirements and/or professional judgment were used as appropriate.

### REVIEW ELEMENTS

The data were evaluated based on the following review elements (where applicable to the method):

- ✓ Data completeness (chain-of-custody [COC])/sample integrity
- ✓ Holding times and sample preservation
- ✓ GC/MS performance checks
- X Initial calibration/continuing calibration verification

- ✓ Laboratory blanks/equipment blanks/trip blanks
- ✓ Surrogate spike recoveries
- NA Matrix spike (MS) and/or matrix spike duplicate (MSD) results
- ✓ Laboratory control sample (LCS) results
- NA Field duplicate results
- ✓ Internal standard results
- ✓ Sample results/reporting issues

The symbol (✓) indicates that no validation qualifiers were applied based on this parameter. NA indicates that the parameter was not included as part of this data set or was not applicable to this validation and therefore not reviewed. The symbol (X) indicates that a QC nonconformance resulted in the qualification of data. Any QC nonconformance that resulted in the qualification of data is discussed below. In addition, nonconformances or other issues that were noted during validation, but did not result in qualification of data, may be discussed for informational purposes only.

The data appear valid as reported and may be used for decision making purposes. Selected data points were estimated due to nonconformances of certain QC criteria (see discussion below). Qualified sample results are presented in Table 1.

## RESULTS

### Data Completeness (COC)/Sample Integrity

The data package was reviewed and found to meet acceptance criteria for completeness:

- The COCs were reviewed for completeness of information relevant to the samples and requested analyses, and for signatures indicating transfer of sample custody.
- The laboratory sample login sheet(s) were reviewed for issues potentially affecting sample integrity, including the condition of sample containers upon receipt at the laboratory.
- Completeness of analyses was verified by comparing the reported results to the COC requests.

### Holding Times/Sample Preservation

Sample preservation and preparation/analysis holding times were reviewed for conformance with the QC acceptance criteria. The QC acceptance criteria were met.

### GC/MS Performance Checks

The data were reviewed to ensure that the 4-bromofluorobenzene (BFB) tuning was performed at the correct frequency and that the method acceptance criteria were met. The QC acceptance criteria were met.

### Initial Calibration/Continuing Calibration Verification

Calibration data were reviewed for conformance with the QC acceptance criteria to ensure that:

- the initial calibration (ICAL) percent relative standard deviation (%RSD), correlation coefficient (r)/coefficient of determination ( $r^2$ ), and/or response factor method acceptance criteria were met;

- the initial calibration verification standard (ICV) percent recoveries (%Rs) acceptance criteria were met;
- the continuing calibration verification standard (CCV) method percent difference or percent drift (%Ds) and RF acceptance criteria were met; and
- the retention time method acceptance criteria were met.

Data qualification to the analytes associated with the specific ICAL and/or CCV was as follows:

#### ICAL Linearity Nonconformances:

Nonconformance	Actions	
	Detected Results	Nondetected Results
%RSD > 15% and quantitation based on mean RF	J	UJ
r or $r^2$ < 0.99 and quantitation based on linear regression	J*	UJ*
* No guidance in NFG, thus RESCON professional judgment was used		

#### CCV Linearity Nonconformances:

Nonconformance	Actions	
	Detected Results	Nondetected Results
%D > 20%	J	UJ
%Drift > 20%	J*	UJ*
* No guidance in NFG, thus professional judgment was used		

Qualified sample results are shown in Table 1. Nonconformances are summarized in Attachment A in Table A-1a and A1b.

#### Laboratory Blanks/Equipment Blanks/Trip Blanks

Laboratory method blanks, equipment rinsate and trip blanks were evaluated as to whether there were contaminants detected above the detection limit (DL).

Data validation qualifications for individual samples are based on the maximum contaminant concentration detected in all associated blanks.

Method, equipment rinsate and trip blank results were reviewed for conformance with the QC acceptance criteria. Detected results in blanks are not discussed in this data validation report if the associated results were nondetect or if qualification of sample results was not required.

The QC acceptance criteria were met and/or qualification of the sample results was not required.

#### Surrogate Spike Recoveries

The surrogate recoveries (%Rs) were reviewed for conformance with the QC acceptance criteria. All QC acceptance criteria were met.

#### MS/MSD Results

MS/MSD analyses were not performed on samples reported in this SDG. There were no validation actions taken on this basis.

### **LCS Results**

The LCS %Rs were reviewed for conformance with the QC acceptance criteria. All QC acceptance criteria were met.

### **Field Duplicate Results**

There were no field duplicate samples submitted with this data set. No validation actions were taken on this basis.

### **Internal Standard Results**

The internal standard (IS) recoveries were reviewed for conformance with the QC acceptance criteria. All QC acceptance criteria were met.

### **Sample Results/Reporting Issues**

Compounds that were not detected in the sample are reported as undetected (U) at the Limit of Detection (LOD).

Compounds detected at concentrations less than the LOQ but greater than the detection limit (DL) were qualified by the laboratory as estimated (J). This "J" qualifier was retained during data validation.

Any sample that was analyzed at a dilution due to high concentrations of target or non-target compounds or matrix interferences was checked to ensure that the results and/or sample specific LODs and LOQs were adjusted accordingly by the laboratory.

### **QUALIFICATION ACTIONS**

Sample results qualified as a result of validation actions are summarized in Table 1. All actions are described above.

### **ATTACHMENTS**

Attachment A: Nonconformance Summary Tables

Attachment B: Qualifier Codes and Explanations

Attachment C: Reason Codes and Explanations

**Table 1 - Data Validation Summary of Qualified Data**

Sample ID	Matrix	Compound	Result	LOD	Units	Validation Qualifiers	Validation Reason
BPOW5-1-GW-061314	WG	4-METHYL-2-PENTANONE		2.5	UG/L	UJ	c
BPOW5-1-GW-061314	WG	CHLOROETHANE		1.0	UG/L	UJ	c
BPOW5-1-GW-061314	WG	DICHLORODIFLUOROMETHANE		1.0	UG/L	UJ	c
BPOW5-1-GW-061314	WG	2-BUTANONE		2.5	UG/L	UJ	c
BPOW5-2-GW-061314	WG	2-BUTANONE		2.5	UG/L	UJ	c
BPOW5-2-GW-061314	WG	4-METHYL-2-PENTANONE		2.5	UG/L	UJ	c
BPOW5-2-GW-061314	WG	CHLOROETHANE		1.0	UG/L	UJ	c
BPOW5-2-GW-061314	WG	DICHLORODIFLUOROMETHANE		1.0	UG/L	UJ	c
BPOW5-3-GW-061314	WG	2-BUTANONE		2.5	UG/L	UJ	c
BPOW5-3-GW-061314	WG	4-METHYL-2-PENTANONE		2.5	UG/L	UJ	c
BPOW5-3-GW-061314	WG	CHLOROETHANE		1.0	UG/L	UJ	c
BPOW5-3-GW-061314	WG	DICHLORODIFLUOROMETHANE		1.0	UG/L	UJ	c
BPOW5-TB-061314	WQ	2-BUTANONE		2.5	UG/L	UJ	c
BPOW5-TB-061314	WQ	4-METHYL-2-PENTANONE		2.5	UG/L	UJ	c
BPOW5-TB-061314	WQ	CHLOROETHANE		1.0	UG/L	UJ	c
BPOW5-TB-061314	WQ	DICHLORODIFLUOROMETHANE		1.0	UG/L	UJ	c

## Attachment A

## Nonconformance Summary Tables

Table A-1a – Initial Calibration

ICAL	Compound	% RSD	Limit
WG144357 GCMS-C	CHLOROETHANE	36.2	< 15%
Associated samples: All samples in the SDG			

Table A-1b - Continuing Calibration Verification

CCV ID	Compound	% D	Limit
WG144955-4 GCMS-C	DICHLORODIFLUOROMETHANE	56.1	< 20%
	2-BUTANONE	20.7	< 20%
	4-METHYL-2-PENTANONE	37.9	< 20%
Associated samples: All samples in this SDG			

**Attachment B****Qualifier Codes and Explanations**

Qualifier	Explanation
J	The analyte was positively identified; the associated numerical value is the approximate concentration of the analyte in the sample.
UJ	The analyte was not detected above the reported sample quantitation limit. However, the reported quantitation limit is approximate and may or may not represent the actual limit of quantitation necessary to accurately and precisely measure the analyte in the sample.
U	The analyte was analyzed for, but was not detected above the reported sample quantitation limit.
R	The sample results are rejected due to serious deficiencies in the ability to analyze the sample and meet quality control criteria. The presence or absence of the analyte cannot be verified.

## Attachment C

## Reason Codes and Explanations

Reason Code	Explanation
be	Equipment blank contamination
bf	Field blank contamination
bl	Laboratory blank contamination
c	Calibration issue
co	Analyte carryover
d	Reporting limit raised due to chromatographic interference
fd	Field duplicate RPDs
h	Holding times
i	Internal standard areas
k	Estimated Maximum Possible Concentration (EMPC)
l	LCS or OPR recoveries
lc	Labeled compound recovery
ld	Laboratory duplicate RPDs
lp	Laboratory control sample/laboratory control sample duplicate RPDs
m	Matrix spike recovery
md	Matrix spike/matrix spike duplicate RPDs
nb	Negative laboratory blank contamination
p	Chemical preservation issue
r	Dual column RPD
q	Quantitation issue
s	Surrogate recovery
su	Ion suppression
t	Temperature preservation issue
x	Percent solids
y	Serial dilution results
z	ICS results
mc	Method compliance nonconformance

